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(54) TILE: PROTEIN HAVING TPO ACTIVITY

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(57) Abstract

The present invention relates to thrombopoletin (TPO) polypeptides having the biological scrivity of specifically stimulating or increasing platelet production compariting the autino acid sequence 1-332 of SEQ ID NO: 6 or a derivative thereof, DNA molecules aconding 1PO polypeptides, processes for production of the polypeptides, ambodies specifically immunoreactive with the polypeptides, pharmaconding compositions compristing the polypeptides, and methods for using the polypeptides in treatment of platelet disorders such as thrombo-trapeals.

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PROTEIN HAVING TPO ACTIVITY

now pending, which is a continuation-in-part of U.S. Patent Application Serial No. 08/221,020 filed April 1, 1994, now abandoned, which is a continuation-in-part of U.S. Patent This application is a continuation-in-part of U.S. Patent Application Serial No. (unknown) filed January 31, 1995, now pending, which is a continuation-in-part of U.S. Patent Application Serial No. 08/361,811 filed December 22, 1994, now pending, which is a continuation-in-part of U.S. Patent Application Serial No. 08/320,300 filed October 11, 1994, now pending, which is a continuation-in-part of U.S. Patent Application Serial No. 08/278,083 filed July 20, 1994, Application Serial No. 08/212,164 filed March 14, 1994 now abandoned.

FIELD OF THE INVENTION

in a specific manner or to enhance proliferation and differentiation of megakaryocyte progenitor cells, to a DNA an activity to stimulate or increase platelet production in vivo sequence coding for said protein, and to a process for the This invention relates to a novel protein which has production of the same.

BACKGROUND OF THE INVENTION

Megakaryocytes · further undergo polyploidization and mainly in bone marrow. Megakaryocytes originate from a nto megakaryocyte progenitor cells, which are committed to the megakaryocytic lineage. Megakaryocyte progenitor cells multinucleate cells which produce platelets and can be found primitive pluripotent stem cell differentiates to some degree proliferate and differentiate into megakaryocytes. cytoplasmic maturation and finally release their anuclear Megakaryocytes are large cytoplasm-rich pluripotent hematopoietic stem cell in bone marrow. A

albuminal surface of the sinus endothellum in the bone marrow Iwo to four thousands of platelets are on an average formed Although the platelet formation mechanism is still unclear in many points, it is considered that megakaryocytes are typically localized on the and produce cytoplasmic processes that extend into the cytoplasmic fragments, namely platelets, into the circulation. sinusoid where they undergo fragmentation of platelets. from one mature megakaryocyte.

layer of the skin of the venous sinus of the bone marrow and that the cytoplasm comes through the dermal skin to result in a cord-like projection on the internal wall of the venous sinus There are many ambiguous points regarding the mechanism for generating the platelets though it is likely that the megakaryocytes are locally present in the dermal whereby the platelets are discharged.

when the numbers of erythrocytes and leucocytes are at a and, finally, they return to the normal level. Also in the clinical field, it has been known that a decrease in the number of the platelets thrombocytopenia and thrombocytosis even It has been suggested that there is a specific function for the production and the control of the megakaryocyte hematopoiesis platelet production. In healthy human beings and animals, the number of the effective platelets is maintained though it has been known that, upon administration of an antiplatelet antibody to the healthy then they begin to increase temporarily to larger than usual animals for example, the platelet number decrease promptly, normal level.

mechanism. If the hemostatic mechanism does not function Incidentally, the most important function of a platelet is the production of a thrombus in a hemostatic properly due to a decrease in the platelet number, a tendency towards hemostasis results.

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and temporarily exceeds the normal level, but finally returns however, success in the isolation and identification of a thrombopoiesis. Platelets are maintained in effective numbers that, when an anti-platelet antibody is administered to a to the normal level. It is known also in the clinical field that a decrease in the number of platelets (thrombocytopenia) or an ncrease in their number (thrombocytosis) occurs even under specific regulatory factor involved in platelet production (for example, like the case of erythropoietin in the erythrocyte The presence of a specific regulatory mechanism has been suggested with regard to megakaryocytopoiesis and in healthy people and normal animals. However, it is known normal animal, only the number of platelets decreases sharply within a short period of time, starts to increase thereafter normal numbers of erythrocytes and leucocytes. To date, formation) has not been reported.

and platelet transfusion is applied to such patients in order to prevent bleeding tendency. Platelet transfusion is also applied Most important function of platelets is the tendency occurs when normal function of the hemostatic radiotherapy and chemotherapy of cancers, thrombocytopenia to patients after bone marrow transplantation or of aplastic Bleeding mechanism is spoiled by thrombocytopenia. In the field of caused by bone marrow suppression is a mortal complication, formation of blood clot in the hemostatic mechanism.

Platelets for use in such platelet transfusion are prepared by plateletpheresis from blood of healthy blood donors, but such platelets for transfusion use have a short bacterial infection. The platelet transfusion also has a possible danger of exposing patients to dangerous viruses such as human immunodeficiency virus (HIV) or various hepatitis shelf life and a possibility of causing contamination by viruses, of inducing antibodies specific for a major

nistocompatibility antigen (HLA) of the transfused platelets or contaminated lymphocytes in the platelets for transfusion use. causing graft versus host disease (GVHD) due

In consequence, it will be of great benefit if intrinsic platelet formation can be stimulated in thrombocytopenic patients and, at the same time, their platelet transfusion dependency can be reduced. In addition, if thrombocytopenia in cancer patients undergoing radiotherapy or chemotherapy can be corrected or prevented, such treatments can be made more safer, the intensity of the reatment can possibly be increased, and further improvement of the anti-cancer effects can be expected.

have been made on the isolation and identification of specific regulatory factors involved in the regulation of megakaryocyte to form megakaryocyte colonies in a semi-solid culture medium. The other regulatory factor, called megakaryocyte actor, thrombopoiesis stimulating factor or the like, acts enhancing their differentiation and maturation. The Meg-Pot is For these reasons, a number of intensive studies Cell. Physiol., vol.110, pp.101 - 104, 1982). Megakaryocyte colony stimulating factor (Meg-CSF) is a regulatory factor potentiating factor (Meg-Pot), megakaryocyte stimulating detectable in combination with Meg-CSF activity in some cases. In addition, since platelet counts increased when serum or plasma collected from experimentally induced and platelet production. According to in vitro studies, regulatory factors governing megakaryocytopoiesis are roughly which stimulates proliferation and differentiation of CFU-MK mainly upon immature or mature megakaryocytes thereby hrombocytopenic animals was administered to other normal divided into the following two factors (cf. Williams $et\ al.,\ J.$ animals, it has been suggested that there is a humoral factor, called thrombopoietin (TPO), capable of promoting platelet

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platelet production. Human IL-6 did not show Meg-CSF nematopoietic cells, it can be distinguished from specific egulatory factors controlling megakaryocytopolesis and activity, but acted upon immature megakaryocytes and then anhanced their differentiation into mature megakaryocytes Williams et al., Exp. Hematol., vol.18, p.69, 1990). In vivo promoted both maturation and shifted to higher ploidy of bone marrow megakaryocytes in primates, but also caused side effects such as reduction of body weight, induction of acute Human IL-11 showed no Meg-CSF activity, but exerted Meg-Pot been cloned were examined for their abilities to stimulate stimulated formation of human megakaryocyte colonies (Bruno exerts its effects upon proliferation and differentiation of all administration of IL-6 induced platelet production and phase protein (Asano et al., Blood, vol.75, pp.1602 - 1605, .990; Stahl et al., Blood, vol.78, pp.1467 - 1475, 1991). In recent years, some cytokines whose genes have et al., Exp. Hematol., vol.16, pp.371 - 377, 1988) and, at least vol.241, p.1820, 1988). However, since IL-3 is a factor which n monkey, increased platelet counts (Donahue et al., Science, megakaryocytopoiesis and thrombopoiesis.

Although clinical application of these cytokines as platelet increasing factors is expected, their functions are not specific for the megakaryocyte lineage and they cause side effects. Therefore, development of a platelet increasing factor which is specific for the megakaryocyte-platelet system and causes less side effects has been called for in the clinical field.

Blood, vol.81, pp.901 - 908, 1993). In addition, human LIF significantly increased platelet counts in primates (Mayer et al., Blood, vol.81, pp.3226 - 3233, 1993), but its in vitro action

Jpon megakaryocytes was weak (Burstein *et al., J. Cell.*

Physiol., vol.153, pp.305 - 312, 1992).

activity and promoted platelet production in mice (Neben et $al.,\,$

Meg-CSF, Meg-pot or TPO activity has been found in serum, plasma or urine of thrombocytopenic patients or animals, or in culture supernatant of certain human cultured cell lines. However, whether these activities are due to the presence of a single factor or a combination of several factors or whether they are different from known cytokines is presently unknown.

found by further studies that the material was not at a level of selenomethionine into newly forming platelets in mice has been partially purified from plasma of thrombocytopenic patients or urine of patients with idiopathic thrombocytopenic patients with hypomegakaryocytic thrombocytopenia (Hoffman et al., J. Clin. Invest., vol.75, pp.1174 - 1182, 1985), but it was Blood, vol.74, pp.1196 - 1212, 1989). A substance having TPOike activity which enhanced incorporation of 75Se-Hematol., vol.12, pp.624 - 628, 1984; de Alarcon and nolecular weight of 46,000 was purified from plasma of surity to allow accurate amino acid sequencing (Hoffman, urpura (ITP), and apparent molecular weight of the plasmaserived factor was determined to be 40,000 (Grossi et al., Hoffman et al. have found that sera of patients with aplastic anemia and amegakaryocytic thrombocytopenic purpura contained a Meg-CSF activity that significantly augmented human megakaryocyte colony formation (Hoffman et al., N. Eng. J. Med., vol.305, pp.533 - 538, 1981). Thereafter, Mazur et al. have reported that the Meg-CSF activity present in he serum of aplastic anemia patients was distinct from both IL-3 and GM-CSF (Mazur et al., Blood, vol.76, pp.290 - 297, 1990). Similar Meg-CSF activity has been found in sera of cancer patients receiving intensive cytotoxic chemotherapy and bone marrow-transplant patients (Mazur et al., Exp. Schmieder, Prog. Clin. Bio. Res., vol.215, pp.335 - 340, 1986). Hoffman et al. have reported that Meg-CSF with an apparent

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urinary samples, but with no information on its structure Erikson-Miller et al., "Blood Cell Growth Factors: their present AiphaMed Press, Dayton, Ohio, pp.204 - 220, 1992). Turner et vol.78, p.1106 279a, 1991, (abstr.,supple. 1)). This MSF has a molecular weight of 28,000 to 35,000. Identity of this factor with the Meg-CSF so far detected in serum and plasma samples of thrombocytopenic patients and its platelet increasing Kawakita et al. have further reported that the Meg-CSF activity found in the urine extract of aplastic anemia patients showed an apparent molecular weight of 45,000 by gel nave also reported on the purification of Meg-CSF from similar and future use in hematology and oncology" ed. by Murphy, al. have purified a megakaryocyte stimulating factor (MSF) having Meg-CSF activity from the urine of bone marrow Meg-CSF and TPO-like activities have also been detected in urine samples of patients with aplastic anemia and severe ITP (Kawakita et al., Br. J. Haematol., vol.48, pp.609 iltration under a dissociation condition (Kawakita et al., Br. J. Haematol., vol.62, pp.715 - 722, 1986). Erikson-Miller et al. transplant patients and cloned its gene (Turner et al., Blood, 515, 1981; Kawakita et al., Blood, vol.556 - 560, 1983). activity remain to be elucidated.

A substance having a TPO-like activity with a molecular weight of 32,000 has been purified from the culture supernatant of a human embryonal kidney-derived cell line (HEK cells) and its biological and biochemical properties have extensively been examined, but its structure is still unknown (McDonald *et al., J. Lab. Clin. Med.*, vol.106, pp.162 - 174, 1985; McDonald, *Int. J. Cell Cloning*, vol.7, pp.139 - 155, 1989). On the contrary, other researchers have reported that the main activity in the conditioned medium of the HEK cells, which enhances megakaryocyte maturation *in vitro*, is due to known

cytokines, namely IL-6 and EPO (Withy et al., J. Cell. Physiol., vol.15, pp.362 - 372, 1992).

and Hill and Levin have conducted partial purification of a TPO-like activity from plasma of thrombocytopenic rabbits nave reported that a TPO-like activity which enhanced platelets in rabbits and mice was found in plasma of thrombocytopenic rabbits induced by antiplatelet serum njection (Evatt et al., J. Lab. Clin. Med.; vol.83, pp.364 - 371, 974). In addition to this, a number of similar studies have seen reported from the 1960's to 1970's (for example, Odell et al., Proc. Soc. Biol. Med., vol.108, pp.428 - 431, 1961; Evatt and Am. J. Physiol., vol.218, pp.1376 - 1380, 1970; Shreiner and evin, J. Clin. Invest., vol.49, pp.1709 - 1713, 1970; and Penington, Br. Med. J., vol.1, pp.606 - 608, 1070). Evatt et al., Evatt et al., Blood, vol.54, pp.377 - 388, 1979; Hill and Levin, Exp. Hematol., vol.14, pp.752 - 759, 1986). Thereafter, purification of this factor was continued, monitoring an activity to enhance differentiation and maturation of megakaryocytes in vitro, namely Meg-Pot activity, to reveal that this activity has an apparent molecular weight of 40,000 to 46,000 when determined by gel filtration (Keller et al., Exp. Hematol., vol.16, pp.262 - 267, 1988; Hill et al., Exp. Hematol., vol.20, pp.354 - 360, 1992). Since IL-6 activity was not detectable in plasma of rabbits with severe acute Serum administration, it was suggested that this TPO-like activity ncorporation of ⁷⁵Se-selenomethionine into newly forming evin, J. Clin. Invest., vol.48, pp.1615 - 1626, 1969; Harker, With regard to animal-derived factors, Evatt et al. vould be due to a factor other than IL-6 (Hill et al., Blood, hrombocytopenia induced by antiplatelet /ol.80, pp.346 - 351, 1992).

Tayrien and Rosenberg have also purified a factor naving an apparent molecular weight of 15,000 from thrombocytopenic rabbit plasma and culture supernatant of

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a rat megakaryocytic cell line, but with no information on its HEK cells, which stimulates production of platelet factor 4 in structure (Tayrien and Rosenberg, J. Biol. Chem., vol.262, op.3262 - 3268, 1987).

In addition, Nakeff has found a Meg-CSF activity in administration (Nakeff, "Experimental Hematology Today" ed. On the other hand, sera from thrombocytopenic rabbits vol.69, pp.1046 - 1052, 1989), but had no detectable Meg-CSF serum of thrombocytopenic mice induced by anti-platelet by Baum and Ledney, Springer-Verlag, NY, pp.111 - 123, 1977). vol.17, pp.903 - 907, 1989) and stimulated morphological enhanced maturation of megakaryocytes (Keller et al., Exp. Hematol., vol.16, pp.262 - 267, 1988; Hill et al., Exp. Hematol., change of megakaryocytes into platelets (Leven and Yee, Blood, activity.

plasma of rats rendered thrombocytopenic by sublethal whole 1984), and suggested that induction of the Meg-CSF activity in latelets, because this activity did not change by platelet 988). Mazur and South have detected a Meg-CSF activity in serum of sublethally irradiated dogs and reported that this actor has an apparent molecular weight of 175,000 when Miura et al. have detected a Meg-CSF activity in vivo is related to decreased megakaryocytes, but not decreased measured by gel filtration (Mazur and South, Exp. Hematol., serum-, plasma- and urine-derived factors have been reported body irradiation (Miura et al., Blood, vol.63, pp.1060 - 1066, ransfusion (Miura et al., Exp. Hematol., vol.16, pp.139 - 144, vol.13, pp.1164 - 1172, 1985). In addition to the above, by other investigators for example by Straneva et al. (Straneva et al., Exp. Hematol., vol.15, pp.657 - 663, 1987).

stimulate megakaryocytopoiesis and thrombopoiesis have been found in biological samples derived from thrombocytopenic Thus, as described above, various activities that patients and animals, but isolation of these factors, their

biochemical and biological identification and determination of their characteristics have not been achieved because of their extremely small contents in natural sources such as blood and

SUMMARY OF THE INVENTION

Objects of the present invention are to isolate a rPO protein from natural sources and identify it, the TPO protein having an activity to stimulate or increase platelet production in vivo and/or to enhance proliferation and the TPO protein and to provide a process for the production of said protein in a homogeneous quality and in a large quantity with recombinant DNA techniques. Success in accomplishing reduction of the currently used platelet transfusion, and such a novel protein will also be used for treatment and diagnosis of referred to as "TPO activity"); and to isolate a gene coding for such objects will lead to the substitution or frequencydifferentiation of megakaryocyte progenitor cells (hereinafter, platelet disorders.

The present invention, therefore, relates to (i) a purified and isolated DNA sequence encoding a protein having TPO activity, which is selected from the group consisting of:

- (a) the DNA sequences shown in SEQ ID NOs: 194, 195 and 196, or complementary strands thereof; and
- (b) DNA sequences which hybridize under stringent conditions to the DNA sequences as defined in (a) or fragments thereof; and
- (c) DNA sequences which would hybridize to the DNA sequences as defined in (a) and (b), but for the degeneracy of the genetic code,
- to a process for the production of a protein having TPO activity, which comprises the steps

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growing, under suitable nutrient conditions, procaryotic or eucaryotic host cells transformed or transfected with said DNA sequence in a manner enabling expression of the protein; and

isolating the desired protein product obtained by expression of said DNA sequence, and

(iii) to a protein product obtained by expression in a procaryotic or eucaryotic host cell of said DNA sequence.

The present invention further relates to a pharmaceutical composition comprising an effective amount of said protein having TPO activity, and to a method for treating platelet disorders, especially thrombocytopenia, comprising administering said protein to patients having the disorders.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 shows Sephacryl S-200HR gel filtration chromatography of Phenyl Sepharose 6 FF/LS F2 derived from XRP.

Figure 2 shows Capcell Pak C1 reverse phase chromatography of YMC-pack CN-AP TPO-active fraction derived from low molecular TPO sample (Sephacryl S-200HR F3) of XRP.

Figure 3 shows a SDS-PAGE analysis of Capcell Pak C1 TPO-active fraction (FA) from low molecular TPO sample of XRP.

Figure 4 shows peptide maps on C18 reverse phase HPLC of rat TPO isolated by SDS-PAGE. The peptide fragments were obtained by systematic hydrolysis with three proteases. Figure 5 shows TPO activity derived from XRP in

the rat CFU-MK assay system. Figure 6 shows a construction of expression vector

PEF18S.

Figure 7 shows TPO activity in the culture supernatant of COS1 cells into which pEF18S-A2 α was introduced in the rat CFU-MK assay system.

Figure 8 shows TPO activity in the culture supernatant of COS1 cells into which pEF18S-HL34 was introduced in the rat CFU-MK assay system.

Figure 9 shows TPO activity in the culture supernatant of COS1 cells into which pHT1-231 was introduced in the rat CFU-MK assay system.

Figure 10a shows TPO activity in the culture supernatant of COS1 cells into which pHTF1 was introduced in the rat CFU-MK assay system.

Figure 10b shows TPO activity in the culture supernatant of COS1 cells into which pHTF1 was introduced in the M-07e assay system.

Figure 11 shows a restriction map of the λHGT1 clone and construction of pHGT1 and pEFHGTE.

(E: EcoRI, H: HindIII, S: Sall)

Figure 12a shows TPO activity in the culture supernatant of COS1 cells into which pEFHGTE was introduced in the rat CFU-MK assay system.

Figure 12b shows TPO activity in the culture supernatant of COS1 cells into which pEFHGTE was introduced in the M-07e assay system.

Figure 13a shows TPO activity in the culture supernatant of COS1 cells into which pHT1-211#1, pHT1-191#1, or pHT1-171#2 was introduced in the rat CFU-MK assay system.

Figure 13b shows TPO activity in the culture supernatant of COS1 cells into which pHT1-163#2 was introduced in the rat CFU-MK assay system.

Figure 14 shows TPO activity in the culture supernatant of COS1 cells into which pHT1-211#1, pHT1-

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191#1, pHT1-171#2, or pHT1-163#2 was introduced in the M-07e assay system.

pDEF202-hTPO-P1 has been introduced and allowed the TPO to Figure 15 is a chromatogram of a reverse phase chromatography (Vydac C4 column) for purification of human TPO from the culture supernatant of CHO cells into which be expressed.

Figure 16 is a photograph showing the SDS-PAGE separation of the human TPO purified from the culture supernatant of CHO cells into which pDEF202-hTPO-P1 has been introduced and allowed the TPO to be expressed.

Figure 17 is a chromatogram of a reverse phase chromatography for purification of human TPO from E. coli into which pCFM536/h6T(1-163) has been introduced and allowed the TPO to be expressed.

Figure 18 is a photograph showing the SDS-PAGE separation of the variant human TPO, h6T(1-163) isolated and purified from E. coli into which pCFM536/h6T(1-163) has been introduced and allowed the TPO to be expressed.

prepared by transfection of the human TPO expression plasmid Figure 19 shows an elution pattern of hTPO163 on Superdex 75 pg column with regard to the purification of the pDEF202-hTPO163 into CHO cells. The protein amount was hTPO 163 from the culture supernatant, as a starting material, determined at 220 nm.

standard hTPO163 eluted from Superdex 75 pg column upon the purification of the hTPO163 from the culture supernatant, as a stating material, prepared by transfection of the human TPO expression plasmid pDEF202-hTPO163 into CHO cells. The Figure 20 shows an SDS-PAGE analysis of the TPO163 was silver-stained on the gel.

Figure 21 shows a structure of the expression vector pSMT201.

as determined by M-07e assay, in the culture supernatant of Figure 22 shows a graph showing the TPO activity the COS7 that BGL-TPO, N3/TPO or 09/TPO has been introduced into and then expressed.

cultures that an insertion or deletion derivative of human TPO Figure 23 is a graph showing a TPO activity, as determined by M-07e assay, in the supernatants of COS7 cell has been introduced into and expressed.

Figure 24 shows increased platelet count in mice administered TPO via intravenous and subcutaneus injection.

Figure 25 shows dose dependent increased platelet count in mice following subcutaneous injection of TPO.

Figure 26 shows TPO induced increase in platelet count following treatment of mice with 5-FU to induce thrombocytopenia.

Figure 27 shows TPO induced increase in platelet nimustine count following treatment of mice with hydrochloride to induce thrombocytopenia.

Figure 28 shows TPO induced increase in platelet count thrombocytopenic mice following bone marrow ransplant.

Figure 29 shows TPO induced platelet count \$ mice irradiation of following x-ray hrombocytopenia.

Figure 30 shows dose-dependent increase in platelet count following administration of truncated TPO (amino acids 1-163 in SEQ ID NO: 6).

Figure 31 shows increase in platelet count following administration of truncated TPO (amino acids 1-163 in SEQ ID NO: 6) following administration of nimustine nydrochloride to induce thrombocytopenia.

Figure 32 shows that increasing concentrations of Mpl-X added to the human megakaryocyte culture system ncreasingly block megakaryocyte development. - 16

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Figure 33 describes TPO activity, as determined by Aig133]TPO(1-163), [Met-2, Lys-1, Ala1, Vaß, Pro148]TPO(1-163) and [Met-2, Lys-1, Ala1, Val3, Arg115]TPO(1-163) M-07e assay, of TPO derivatives [Met-2, Lys-1, Ala1, Val3, expressed in E. coli.

Alg129JTPO(1-163), [Met-2, Lys-1, Ala1; Val3, Arg143JTPO(1-163), [Met-2, Lys-1, Ala1, Val3, Leu82]TPO(1-163), [Met-2, Lys-Figure 34 describes TPO activity, as determined by 1, Ala1, Val3, Leu146]TPO(1-163) and [Met-2, Lys-1, Ala1, Val3, M-07e assay, of TPO derivatives [Met-2, Lys-1, Ala1, Val3, Arg⁵⁹]TPO(1-163) expressed in E. coli.

DETAILED DESCRIPTION OF THE INVENTION

polypeptides of the invention include [Thr³³, Thr³³³, Ser³³⁴, IIe³³⁵, Gly³³⁶, Tyr³³⁷, Pro³³⁸, Tyr³³⁹, Asp³⁴⁰, Val³⁴¹, Pro³⁴², Asp³⁴³, Tyr³⁴⁴, Ala³⁴⁵, Gly³⁴⁶, Val³⁴⁷, His³⁴⁸, His³⁴⁹, His³⁵⁰, His³⁵³, His³⁵³, Thr³³³, Ser³³⁴, IIe³³⁵, Gly³³⁶, Tyr³³⁷, Pro³³⁸, Tyr³³⁹, Asp³⁴⁰, Val³⁴⁷, His³⁴⁸, His³⁴⁸, His³⁵⁹, His³⁵⁹ invention include derivatives of a TPO polypeptide to 151 of SEQ ID NO: 6; the mature sequence of amino acids of SEQ ID NOs: 2, 4 and 6; including those having from 1 to 6 NH $_2$ Asn²⁵JTPO and [Thr³³JTPO. Other polypeptides of the (AHis³³]TPO(1-163), [AArg¹¹⁷]TPO(1-163), [AGIy¹¹⁶]TPO(1nvention are thrombopoletin (TPO) polypeptides having the to 332 of SEQ ID NO: 6 or derivatives thereof. Illustrative polypeptides include those which consist of the amino acid Additional illustrative Specifically provided according to the present biological activity of specifically stimulating or increasing platelet production and comprising the amino acid sequence 1 sequences 1 to 163 of SEQ ID NO: 6; 1 to 232 of SEQ ID NO: 6; 1 163), [His³³, Thr³³', Pro³⁴]TPO(1-163), terminal amino acids deleted.

[Ala1, Val3, Arg129]TPO(1-163), [Ala1, Val3, Arg133]TPO(1-Arg¹¹⁷]TPO(1-163), [Gly¹¹⁶, Gly¹¹⁶', Arg¹¹⁷]TPO(1-163), Leu82JTPO(1-163), [Ala¹, Val³, Leu¹46JTPO(1-163), [Ala¹, Val³, 21034]TPO(1-163), [His33, Gly33', Pro34, Ser38]TPO(1-163) 163), [Ala¹, Val³, Arg¹43]TPO(1-163), [Ala¹, Val³, Pro148]TPO(1-163), [Ala¹, Val³, Arg⁵⁹]TPO(1-163), and [Ala¹, Gly¹¹⁶, Asn¹¹⁶', Arg¹¹⁷JTPO(1-163), [Gly¹¹⁶, Ala¹¹⁶' /al3, Arg115]TPO(1-163).

may further comprise the amino acids [Met-2-Lys-1], [Met-1] or [Gly-1]. DNAs provided by the invention include those which encode the TPO polypeptide and derivatives described above and are suitably provided as cDNA, genomic DNA and The TPO polypeptides of the invention also include those which are covalently bonded to a polymer, preferably polyethylene glycol. The TPO polypeptides of the invention manufactured DNAs.

producing a TPO polypeptide as described above comprising the steps of expressing a polypeptide encoded by a DNA of the invention in a suitable host, and isolating said TPO polypeptide. Where the TPO polypeptide expressed is a Met-2-Lys-1 polypeptide, such processes can further include the step Also provided by the invention are processes for of cleaving Met-2-Lys-1 from said isolated TPO polypeptide.

GST polypeptide, a thrombin recognition peptide and a TPO Also provided by the invention are processes for producing TPO polypeptides such as glutathione-S-transferase GST) fusion polypeptides. DNA encoding an amino terminal polypeptide is introduced into a suitable host, the fusion polypeptide is isolated and the GST moiety removed by reatment with thrombin. Resulting TPO polypeptides are of the [Gly-1] structure.

Additionally provided are procaryotic or eucaryotic host cells transformed or transfected with a DNA sequence according to the invention in a manner enabling said host cell

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to express a polypeptide having the biological activity of specifically stimulating or increasing platelet production.

Pharmaceutical compositions of the invention comprise an effective amount of a TPO polypeptide or derivative in combination with a pharmaceutically acceptable carrier and are susceptible to use in treatment of platelet disorders, particularly the treatment of thrombocytopenia, such as is induced by chemotherapy, radiotherapy or bone marrow transplantation. Corresponding treatment methods are provided by the invention.

Finally, the present invention provides antibodies specifically immunoreactive with TPO polypeptides and derivatives thereof as described above. Such antibodies are useful in methods for isolation and quantification of TPO polypeptides of the invention.

According to the present invention, there is provided a novel DNA sequence which encodes a protein having a TPO activity (referred to as "DNA sequence of the present invention", hereinafter). The DNA sequence of the present invention includes a DNA sequence which encodes the amino acid sequence shown in SEQ ID NO: 2, 4 or 6 of the Sequence Listing attached hereto.

Also included in the DNA sequence of the present invention is a DNA sequence which encodes a partly modified (substitution, deletion, insertion or addition) version of the aforementioned amino acid sequence shown in SEQ ID NO: 2, 4 or 6, provided that such modifications do not spoil the TPO activity. That is, DNA sequences coding for TPO derivatives are also included in the present invention.

In other words, the DNA sequence of the present invention includes DNA sequences which encode protein molecules whose amino acid sequences are substantially the amino acid sequences shown in SEQ ID NO: 2, 4 or 6. The wording "amino acid sequences are substantially the amino

acid sequences shown in SEQ ID NO: 2, 4 or 6" as used herein means that said amino acid sequences include those represented by SEQ ID NO: 2, 4 or 6 as well as those represented SEQ ID NO: 2, 4 or 6 which have partial modification therein such as substitution, deletion, insertion, addition or the like, provided that such modifications do not spoil the TPO activity.

Further, the DNA sequence of the present invention consists essentially of a DNA sequence encoding a protein having a TPO activity.

The wording "a DNA sequence which encodes the amino acid sequence" includes all the DNA sequences which may have degeneracy in nucleotide sequences.

· The DNA sequence of the present invention also comprises the following sequences:

(a) DNA sequences represented by SEQ ID No: 7, 194, 195 and 196, or complementary strands thereof,

(b) DNA sequences which hybridize under stringent conditions, to the DNA sequences as defined in (a) or fragments thereof, or

(c) DNA sequences which would hybridize to the DNA sequences as defined in (a) and (b), but for the degeneracy of the genetic code.

In other words, the DNA sequence of the present invention also comprises the following sequences:

(a) DNA sequence which is integrated in the Vector pEF18S-A2α (deposit No. FERM BP-4565) carried by an *E. coli* strain DH5, a Vector pHT1-231 (deposit No. FERM BP-4564) carried by an *E. coli* strain DH5, a Vector pHTF1 (deposit No. FERM BP-4617) carried by an *E. coli* strain DH5, a Vector pHGT1 (deposit No. FERM BP-4616) carried by an *E. coli* strain DH5, and encodes the amino acid sequence of a protein having the TPO activity: or

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stringent conditions) to the DNA sequences defined in (a) or ragments thereof, and encode the amino acid sequence of a hybridize (under (b) DNA sequences which protein having the TPO activity.

ybridization conditions are those employed in the examples degenerate and/or unique sequence oligonucleotide primers See also for example, Chapters 11 and 14 of Laboratory Press, 1989) and Unit 2.10 in Current Protocols in addressing PCR amplification of DNAs of the invention using Molecular Cloning (Sambrook et al., Cold Spring Harbor Motecular Biology, Ausubel et al., Eds., Current Protocols, USA, As used herein, representative "stringent" probes). 1993).

invention is a DNA sequence which encodes a protein having TPO activity and comprises the nucleotide sequence encoding positions 1-163 of the amino acid sequence represented by SEQ Also included in the DNA sequence of the present ID NO: 6.

Such DNA sequences may be supplemented with a restriction enzyme cleavage site and/or an additional DNA sequence at the initiation, termination or intermediate site which facilitate construction of readily expressed vectors. When a non-mammalian host is used, a preferred codon for the gene expression in the host may be incorporated.

An example of the DNA sequence of the present invention is a cDNA molecule which is obtained by preparing mRNA from cells of mammals, including human, and then screening the cDNA in the usual way from a cDNA library prepared by a known method. Sources of the mRNA of this case nclude cells of a rat hepatocyte-derived cell line McA-RH8994, HTC cells, H4-II-E cells, rat liver, kidney, brain and small intestine human liver and the like.

present invention is a genomic DNA molecule which is obtained Another example of the DNA sequence of the

prepared by a known method from cells of mammals including human. Sources of the genomic DNA of this case include by screening it in the usual way from a genomic library chromosomal DNA preparations obtained from human, rat, mouse and the like.

A DNA sequence which encodes a TPO derivative encodes a protein having TPO activity, by modifying the cDNA sequence making use of the known site-directed mutagenesis thereby effecting partial modification of the corresponding may be obtained from the thus obtained cDNA sequence that amino acid sequence.

Having elucidated the amino acid sequence or DNA sequence of a protein having TPO activity in the present invention, a DNA sequence which encodes a partially modified amino acid sequence can be obtained easily by chemical synthesis.

The DNA sequence of the present invention is a useful material for the large scale production of a protein naving TPO activity making use of various recombinant DNA techniques.

species which can be used as a eucaryotic host for large scale DNA coding for TPO of other mammalian species. It is also useful in the gene therapy of human and other mammalian species. In addition, the DNA sequence of the present invention is useful for the development of a transgenic mammalian Also, the DNA sequence of the present invention is useful as a labeled probe for the isolation of a gene which encodes a TPO-related protein, as well as cDNA and genomic production of TPO (Palmiter et al., Science, vol.222, pp.809 814, 1983).

said vector and a process for the production of a protein having in which the aforementioned DNA sequence coding for a protein having TPO activity is integrated, host cells transformed with Also provided by the present invention are a vector

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separating and purifying the expressed protein having TPO

activity.

yeasts include, a baker's yeast (Saccharomyces cerevisiae), a cells of prokaryotes, such as, E. coli and the like, and of Chinese hamster ovary (CHO) cells, C-127 cells, baby hamster kidney (BHK) cells and the like. Illustrative examples of Examples of host cells useful in this case include eukaryotes, such as yeasts, insects, mammals and the like. Ilustrative examples of mammalian cells include COS cells, methanol assimilating yeast (Pichia pastoris) and the like. Ilustrative examples of insect cells include, silkworm cultured cells and the like.

recombinant virus construction, for example, pAc373 (Luckow With regard to vectors for use in the 8, 466 - 472, 1988) or the like may be used. As for yeast cells, pG-1 (Schena M. and Yamamoto K.R., Science, 241, 965 -967, 1988) or the like may be used. A transfer vector for et al., Bio/Technology, 6, 47 - 55, 1988) may be used for the transformation of these host cells, pKC30 (Shimatake H. and M. Rosenberg, Nature, 292, 128 - 132, 1981), pTrc99A (Amann E. et al., Gene, 69, 301 - 315, 1988) or the like may be used for the transformation of E. coli cells. For transforming Genet., 1, 327 - 341, 1982), pCAGGS (Niwa et al., Gene, 108, mammalian cells, pSV2-neo (Southern and Berg, J. Mol. Appl. 193 - 200, 1991), pcDL-SRa296 (Takebe et al., Mol. Cell. Biol., ransformation of silkworm cells.

and the like, as well as an RNA splice site, a polyadenylation signal and the like in the case of vectors for use in eukaryotic contain a replication origin, selection marker(s), a promoter As occasion demands, each of these vectors may

With regard to the replication origin, a sequence derived from, for example, SV40, adenovirus, bovine papilloma

A sequence derived from CoIE1, R factor, F factor or the like from 2 µm DNA, ARS1 or the like may be used in vectors for may be used in vectors for E. coli cells. A sequence derived virus or the like may be used in vectors for mammalian cells.

reast cells. A promoter derived from a nuclear polyhedrosis adenovirus, SV40 and the like may be used in vectors for AOX1 promoter or the like in vectors for methanol assimilating As to promoters for gene expression, those which ior example, trp, Ipp, Iac or tac promoter, may be used in vectors for E. coli cells. ADH, PHO5, GPD, PGK or MAF α promoter may be used in vectors for baker's yeast cells, and are derived from, for example, retrovirus, polyoma virus, mammalian cells. A promoter derived from bacteriophage \(\mathcal{A}_{\tau} \) virus may be used in vectors for silkworm cells.

Typical examples of selection markers useful in resistance gene, a thymidine kinase (TK) gene, a dihydrofolate reductase (DHFR) gene, an E. coli xanthine-guanine Illustrative examples of selection markers useful in vectors for E. coli cells include a kanamycin resistance gene, an ampicillin resistance gene, a tetracycline resistance gene and the like, and those for yeast cells include Leu2, Trp1, Ura3 and phosphoribosyltransferase (${\it Ecogpt}$) gene and the like. vectors for mammalian cells include a neomycin (neo) the like genes.

vector, culturing the resulting transformant and then Production of a protein having TPO activity making use of appropriate combinations of these host-vector systems may be performed by transforming appropriate host cells with a recombinant DNA obtained by inserting the gene of the present invention in an appropriate site of the aforementioned separating and purifying the polypeptide of interest from the esulting cells, or culture medium or filtrate. Commonly used - 24

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When expressing the gene of interest, the original signal sequence may be modified or replaced by a signal sequence derived from another protein in order to obtain homogeneous N-terminal of the expressed product. Homogenization of the N-terminal may be carried out by modification (substitution or addition) of the amino acid residues in the N-terminal or its vicinity. In the case of expression by using E. coli, as a host cell, for example, lysine residue may be further supplemented in addition to methionine residue.

The novel protein of the present invention having TPO activity (to be referred to as "protein of the present invention" hereinafter) includes proteins each of which comprising the amino acid sequence shown in SEQ ID NO: 2, 4 or 6. TPO derivatives whose amino acid sequences are partially modified (substitution, deletion, insertion or addition) are also included in the present invention, provided that the TPO activity is not spoiled by the modification.

In other words, the protein of the present invention includes protein molecules whose amino acid sequences are substantially the amino acid sequences shown in SEQ ID NO: 2, and 6

The wording "amino acid sequences are substantially the amino acid sequences shown in (or represented by) SEQ ID NO: 2, 4 or 6" as used herein means that said amino acid sequences include those shown in SEQ ID NO: 2, 4 or 6 which have partial modification therein such as substitution, deletion, insertion, addition and the like, provided that such modifications do not spoil the TPO activity.

The protein of this invention includes a protein containing the positions 7-151 of the amino acid sequence

shown in SEQ ID NO: 6 and having a TPO activity. Also included in the protein of the present invention is a protein which has TPO activity and comprises positions 1-163 of the amino acid sequence represented by SEQ ID NO: 6.

Examples of other TPO derivatives of the present invention include a derivative whose stability and durability in vivo were improved by amino acid modification (substitution, deletion, insertion or addition), a derivative in which at least one potential glycosylation was changed by deletion or addition, a derivative in which at least one cysteine residue was deleted or substituted by other amino acid residue (alanine or serine residue for example).

Preferably, the protein of the present invention is characterized in that it is separated and purified from host cells transformed with a recombinant vector containing a cDNA molecule, a genomic DNA molecule or a DNA fragment obtained by chemical synthesis.

When intracellular expression is effected using a bacterium, such as *E. coli* as the host, a protein in which an initiation methionine residue is added to the N-terminal side of a protein molecule having TPO activity is obtained, which is also included in the present invention. Depending on the host used, the produced protein having TPO activity may or may not be glycosylated, and each of such cases is included in the protein of the present invention.

The protein of the present invention also includes naturally occurring TPO-active proteins purified and isolated from natural sources such as cell culture medium having TPO activity or human urine, serum and plasma.

A process for the purification of TPO from such natural sources is also included in the present invention. Such a purification process may performed by employing one of or a combination of usually used protein purification steps such as an ion exchange chromatography, a lectin affinity

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affinity chromatography, a sulfated gel chromatography, a produced by recombinant DNA techniques using CHO cell as a chromatography, a triazine dye affinity chromatography, a lydrophobic interaction chromatography, a gel filtration chromatography, a reverse phase chromatography, a heparin lydroxylapatite chromatography, isoelectric focussing certain techniques are used in combination making use of physicochemical properties of TPO which can be deduced from the Examples of this specification is also included in the In addition, an antibody affinity chromatography to which an antibody capable of recognizing FPO is applied may also be used. Furthermore, it has been ound that TPO is the ligand for Mpl (de Sauvage et al., Nature 369: 533-538 (1994); Bartley et al., Cell 77: 1117-1124 (1994); Kaushansky et al., Nature 369: 565-568 (1994)), by hat the Mpl has been coupled to a resin. More particularly, an example of the column is an Mpl-X column which is prepared by coupling a resin with the extracellular region of Mpl (Mpl-X) preparative electrophoresis, an isoelectric focussing gel electrophoresis and the like. A purification process in which which the TPO may be purified using an affinity gel column chromatography, a metal chelating chromatography, nost (Bartley et al. (1994), <u>supra</u>). present invention.

As disclosed herein, the TPO polypeptides of this nvention are further characterized by the ability to bind to the Mpl receptor and specifically to the extracellular (soluble) domain thereof.

protein-coding strand of a human cDNA or genomic DNA sequence of TPO gene, namely a "complementary inverted Also included in the present invention is a protein encoded by a DNA moiety which is complementary to the protein" disclosed by Tramontano et al. (Nucl. Acids Res., vol.12, pp.5049 - 5059, 1984).

marker, for example, 1251 labeling or biotinylation, thus expressing cells in solid samples such as tissue and liquid of the present invention which is labeled with a detectable rendering possible provision of a reagent which is useful for the detection and quantification of TPO or TPO receptor-Also included in the present invention is a protein samples such as blood, urine and the like.

The biotinylated protein of the present invention is order to remove megakaryoblasts from bone marrow at the time of autogenous bone marrow transplantation. It is also useful in the case of its binding to immobilized streptavidin in order to concentrate autogenous or allogenic megakaryocytic diphtheria toxin or the like or with a radioactive isotope is useful in antitumor therapy and in conditioning for bone useful in the case of its binding to immobilized streptavidin in cells at the time of autogenous or allogenic bone marrow transplantation. A conjugate of TPO with a toxin such as ricin, narrow transplantation.

material which is useful when labeled with a detectable marker including a radioactive marker or a non-radioactive marker such as biotin or when it is used in a hybridization procedure for the detection of the position of human TPO gene and/or its related gene family on a chromosomal map. Such a material is also useful for the confirmation of human TPO gene . The present invention also provides a nucleic acid disorders at the DNA level and can be used as a genetic marker for the confirmation of adjoining genes and their disorders.

effective amount of the protein of the present invention The term "therapeutically effective amount" as used herein neans an amount which provides a therapeutic effect for pharmaceutical composition which contains a therapeutically together with a useful and effective diluent, antiseptic agent, solubilizing agent, emulsifying agent, adjuvant and/or carrier. Also included in the present invention is

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TPO protein) of body weight depending on conditions and sex of The pharmaceutical composition comprising the protein of the present invention may be administrated several limes per day, usually in an amount of 0.05µg to 1mg/kg (as a patient, administration route and the like.

of 25,000-500,000,000 of the active ingredient (in terms of protein of the present invention can be administered at a dose The pharmaceutical composition containing the the relative activity by the M-07e assay which will be given later) per kg of body weight for one to several times a day and for one to seven days a week depending upon the symptoms, sex and route of administration.

152nd amino acid sequence shown in SEQ ID NO: 6 are deleted and that, with respect to the N-terminal site, the activity is The present inventors have confirmed that, with respect to the C-terminal site of the human TPO, the activity is maintained even when the amino acid residues up to the maintained even when the amino acid residues up to the 6th amino acid sequence are deleted.

covalent bonding of the protein of the present invention to a

polymer such as polyethylene glycol, chelation of the protein

with metal ions, incorporation of the protein into a granular

preparation, or on the surface, consisting of a polymer

compound such as polylatic acid, polyglycolic acid or hydrogel,

microemulsion, micelles, single or multilayer vesicles, red cell ghosts or spheroplasts. Such a composition will exert nfluences upon physical conditions, solubility, stability, in

or incorporations of the protein into liposomes,

as lactose or mannitol. Also used is a composition comprising

senzyl alcohol or paraben and a vehicle or tonicity agent such

sodium metabisulfite, an antiseptic agent such as thimerosal,

surface active agent such as Tween 20, Tween 80, Pluronic -68 or bile acid salt, a solubilizing agent such as glycerol or oolyethylene glycol, an antioxidant such as ascorbic acid or

Such a composition is used in the form of liquid, freeze-dried or dried preparation and comprises a diluent selected from various buffers (Tris-HCI, acetate and phosphate for example) having various pH values and ionic strengths, a surface adsorption preventing additive such as albumin or gelatin, a

specified conditions and routes of administering conditioning.

addition) in other parts may be preferably used as the effective ingredient of the present invention as well. More preferred TPO derivative is that having the 1st to 163rd activity, containing the 7th to 151th amino acid sequence of SEQ ID NO: 6 and being modified (substitution, deletion, insertion or Accordingly, protein having TPO amino acid sequence of SEQ ID NO: 6.

> chemical properties of the used TPO-active protein. Also ncluded in the present invention is a granular composition in which granules are coated with a polymer such as poloxamer or ooloxamine and TPO which binds to antibodies for tissue specific receptors, ligands or antigens or to tissue specific eceptor ligands. Other examples of the composition of the present invention are those which are in the form of granules, have a protective coating and contain a protease inhibitor or a penetration enhancer, for use in various routes of administration such as parenteral, pulmonary, transnasal and

he composition may be decided depending on the physical and

vivo releasing rate and in vivo clearance of TPO. Selection of

10, IL-11, IL-12, IL-13, LIF and SCF, in addition to the protein A composition which contains at least one of additional hematopoietic factors such as EPO, G-CSF, GM-CSF, M-CSF, IL-1, IL-2, IL-3, IL-4, IL-5, IL-6, IL-7, IL-8, IL-9, ILof the present invention, is also included in the present invention.

The protein of the present invention is useful for the treatment of various thrombocytopenic diseases, either

oral administration.

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Examples of other additional hematopoietic factors 5, IL-6, IL-7, IL-8, IL-9, IL-10, IL-11, IL-12, IL-13, LIF and nclude EPO, G-CSF, GM-CSF, M-CSF, IL-1, IL-2, IL-3, IL-4, ILin combination with other additional hematopoletic alone or factors.

diminished production of TPO. Thrombocytopenia due to reduction in life of platelets or megakaryocytes includes autotransfusion of platelets, in which TPO is administered to a number of the patient's own platelets and the thus increased olatelets are used as transfusion platelets at the time of the it is also useful in the treatment of thrombocytopenia due to acquired immunodeficiency syndrome (AIDS), disseminated intravascular coagulation syndrome and thrombotic hrombocytopenia. In addition, it is also useful in the case of patient prior to a surgical operation in order to increase the production of platelets or reduction in life of platelets (caused by destruction or consumption of platelets), which can be Fanconi anemia or aplastic anemia caused by chemotherapy or rradiation, myelodysplastic syndrome, acute myelocytic diopathic thrombocytopenic purpura, hypoplastic anemia, There are many diseases characterized by platelet disorders such as thrombocytopenia due to damages in the it can be used for the enhancement of the restoration of reated with the protein of the present invention. For example, platelets in a thrombocytopenic patient due to congenital leukemia, aplastic crisis after bone marrow transplantation. peration.

Other use of the protein of the present invention is platelets caused by other chemical or pharmaceutical drugs or curative treatments. TPO can be used for the purpose of enhancing release of new "intact" platelets reatment of diseases resulting from temporal deletion or n such patients. damage

human TPO itself can be used as the antigen or, alternatively, a analyzing the epitope of the antibody. In such cases, these antibodies each having the thus-clarified epitope can be utilized to fractionate, detect, quantitate and purify various rPOs which differ in their properties such as the kind of the can be used as antigens, and the corresponding antibodies nclude both monoclonal and polyclonal antibodies and chimeric antibodies, namely "recombinant" antibodies, prepared by the epitope can be defined by specifying the antigen region. On the other hand, where an antibody to the antigen protein (TPO) tself is utilized, the antigen region can be clarified by antibody specific for TPO. The protein of the present invention conventional methods. To prepare such an anti-TPO antibody, oartial peptide of human TPO can be utilized as the antigen. Where an antibody to the antigen of such a peptide is utilized, present invention further relates to sugar chains added, the length of the peptide chains, etc.

protein, for example, albumin, KLH (key hole limpethemocyanine) or the like, by covalent bonding to prepare an immunoreactive antigen. Alternatively, a peptide of a antigen along with an adjuvant or the like, immunized are mammals, birds, etc. which are generally utilized to make sheep, chickens, hamsters, horses, guinea pigs, etc. From the hus-immunized animal, recovered was its anti-serum and cells, from which a polyclonal antibody is obtained. Where a specifying the antigen region. In this case, the regions of arious TPO molecules having different molecular weights are A TPO peptide containing the thus-selected amino acid sequence is synthesized, and bonded to a suitable carrier multiple antigen peptide (MAP) type is produced to be an USA) 85:5409-5413, 1988). Then, with the thus-prepared hem produce antibodies, such as rabbits, mice, rats, goats, septide is used as the antigen, the epitope can be defined by antigen, according to the Tam's method (Proc.Natl.Acad.Sci.

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screened and identified by immunological engineering. As one example, a peptide-deleted region can be screened and identified. In addition, an antibody gene or a part of the gene is cloned from the cells expressing the intended antibody to obtain an antibody molecule that has been expressed by genetic engineering.

In comparison with a polyclonal antibody which generally contains various antibodies specific for various antigenic determinants (epitopes), a monoclonal antibody is specific for a single antigenic determinant on the antigen. A TPO-specific antibody is useful in improving selectivity and specificity of an antigen-antibody reaction-aided diagnosis and an analytical assay method and in performing separation and purification of TPO. In addition, such antibodies can be applied to the neutralization or removal of TPO from serum. Monoclonal antibodies are also useful for detection and quantification of TPO in, for example, serum of whole blood.

for purification and isolation of human TPO. To fix the chromatography, any conventional methods for fixing various enzymes can be employed. For instance, employable is a method of using a carrier of CNBr-activated Sepharose 4B (produced by Pharmacia Fine Chemicals Co.) or the like. To the column. By this operation, a large amount of human TPO is elution, for example, usable are glycine-HCI buffer (pH 2.5), sodium chloride solution, propionic acid, dioxane, ethylene The anti-human TPO antibody of the present nvention can be used as the ligand in affinity chromatography antibody so as to make it useful in such affinity actually purify human TPO by the use of the fixed anti-human TPO antibody, the fixed anti-human TPO antibody is filled into a column and a human TPO-containing liquid is passed through adsorbed onto the carrier in the column. As the solvent for glycol, chaotropic salt, guanidine hydrochloride, urea, etc.

the elution with such a solvent, human TPO having a high purity

The antibody of the present invention can be used for determining human TPO by immunochemical quantitative assay, especially by enzyme-immunoassay to be conducted by a solid phase sandwich process.

An advantage of monoclonal antibodies is that they can be produced by hybridoma cells in a medium which does not contain any other immunoglobulin molecules. Monoclonal antibodies can be prepared from culture supernatants of hybridoma cells or from mouse ascites induced by intraperitoneal injection of hybridoma cells. The hybridoma technique originally disclosed by Kohler and Milstein (*Eur. J. Immunol. 6:*511-519, 1976) can be used broadly for the formation of hybrid cell lines which are possessed of high level monoclonal antibodies for a number of specific antigens.

as cation-exchange chromatography, anion-exchange chromatography, lectin affinity chromatography, dye adsorption chromatography, hydrophobic interaction etc.) which are generally employed for purification of protein affinity purification can also be employed, in which a gel carrier or membrane, to which has been chemically bonded a To isolate the intended antibody from the thusanti-immunoglobulin-fixed gel chromatography, etc.; as well chromatography, gel permeation chromatography, reversed an be combined. Apart from these, a method of antigen luoro-apatite chromatography, metal chelating obtained antibody-containing material, such as anti-serum, etc., by, purifying it, one or more steps (affinity chromatography such as protein A affinity chromatography, protein G affinity chromatography, Avid gel chromatography, isoelectric point chromatography, partitioning electrophoresis, isoelectric point electrophoresis, phase chromatography, hydroxyl-apatite chromatography chromatography,

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human TPO protein itself or a peptide that contains the region of the antigen or a part of the region or, that is, a molecule capable of recognizing the intended antibody, is prepared, an antibody-containing material is added thereto so that the intended antibody is made to be adsorbed onto the carrier or membrane, and the thus-adsorbed antibody is then eluted and recovered under suitable conditions.

The present invention also allows use of endogenous DNA sequences encoding a TPO polypeptide for the production of large quantities of polypeptide products. For example, *in vitro* and *ex vivo* homologous recombination procedures may be employed to transform host cells to provide for expression or enhanced expression of polypeptides. Preferred host cells include human cells (*e.g.*, liver, bone marrow and the like) wherein a promoter or enhancer sequence is inserted employing flanking sequences that are homologous to a target region in a cellular genome with the result that TPO polypeptide expression is accomplished or enhanced. See, *e.g.*, U.S. Letters Patent 5,272,071, published PCT WO 90/14092, WO 91/06666 and WO 91/09955.

Also provided by the invention are processes for producing a TPO polypeptide as described above comprising the steps of expressing a polypeptide encoded by a DNA of the invention in a suitable host, and isolating said TPO polypeptide expressed is a Met-2-tys-1 polypeptide, such processes can further include the step of cleaving Met-2-tys-1 from said isolated TPO polypeptide.

Also provided are methods for producing TPO polypeptides having [Gly-1] structure comprising the steps of introducing into a suitable host cell a DNA encoding a glutathione-S-transferase (GST) polypeptide 5' to TPO polypeptide encoding sequences, the GST and TPO polypeptide encoding sequences separated by DNA encoding a thrombin recognition polypeptide, isolating the GST-TPO expression

product and treating the expressed polypeptide with thrombin to remove GST amino acids. The resulting TPO polypeptides have a [Gly-1] structure.

The following describes the present invention in

detail.

(A) Purification of rat TPO, analysis of partial amino acid sequences of purified rat TPO and analysis of biological characteristics of purified rat TPO

The inventors of the present invention have firstly attempted to purify a protein (rat TPO) which has an activity to enhance proliferation and differentiation of rat CFU-MK. In this purification study, many trial and error attempts were made on the purification procedures, such as selection of various natural supply sources, selection of gels for chromatographies and separation modes. As the result, the present inventors have succeeded in purifying a protein having a TPO activity from blood plasma of thrombocytopenic rats induced by X-ray or γ -ray irradiation, making use of the TPO activity as a marker based on a rat CFU-MK assay which will be described later in <Reference Example>, and in determining partial amino acid sequences of the purified protein <Examples 1 and 2>.

Biological characteristics of the plasma-derived rat TPO were also examined < Example 3>.

An outline of the procedures from the purification of rat TPO to the determination of partial amino acid sequences of the purified protein is shown in the following.

(i) A blood plasma sample was prepared from about 1,100 thrombocytopenic rats induced by X-ray or γ -ray irradiation and subjected to a Sephadex G-25 chromatography, an anion exchange chromatography (Q-Sepharose FF) and a

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- (ii) Next, the thus obtained WGA-Agarose-adsorbed TPO-active fraction was subjected to a triazine dye affinity chromatography (TSK AF-BLUE 650MH), a hydrophobic interaction chromatography (Phenyl Sepharose 6 FF/LS) and a gel filtration chromatography (Sephacryl S-200 HR) in that order. Since the TPO activity was divided into 4 peaks (F1 as the highest molecular weight fraction, followed by F2, F3 and F4) by the Sephacryl S-200 HR gel filtration, each of the TPO-active fractions F2 and F3 was concentrated to obtain a high molecular weight TPO sample F2 and a low molecular weight TPO sample F3 which were used separately in the subsequent purification steps.
 - (iii) The low molecular weight TPO sample F3 was subjected to a preparative reverse phase chromatography (YMC-Pack PROTEIN-RP), a reverse phase chromatography (YMC-Pack CN-AP) and another reverse phase chromatography (Capcell Pack C1) in that order. When the thus obtained TPO-active fraction was applied to Sodium dodecylsulfate (SDS) polyacrylamide gel electrophoresis (SDS-PAGE) and the TPO-active substance was extracted from the gel, the presence of the TPO activity was confirmed in a band corresponding to apparent molecular weights of about 17,000 to 19,000 under a non-reducing condition.
- applied to SDS-PAGE under a non-reducing condition and transferred on a PVDF membrane. By carrying out systematic limited enzymatic hydrolysis of the protein on the PVDF membrane into peptide fragments, partial amino acid sequences of the rat TPO protein were determined. Based on the amino acid sequence information of two peptide fragments, cloning of rat TPO gene was performed.

(v) Separately from this, the high molecular weight TPO sample F2 obtained by the Sephacryl S-200 HR was subjected to purification in the same manner as in the case of the low molecular weight TPO sample F3. When a TPO-active fraction obtained by the final step reverse phase chromatography (Capcell Pack C1) was applied to SDS-PAGE and the TPO-active substance was extracted from the gel, the presence of the TPO activity was confirmed in a band corresponding to apparent molecular weights of about 17,000 to 22,000 under a non-reducing condition.

(B) Specialization of rat TPO producing cells, preparation of mRNA and construction of rat cDNA library

Since the thus purified rat TPO has been derived from blood plasma, it was necessary to screen organs or cells as the source of mRNA for use in the cDNA cloning. In consequence, TPO activities in various organs and cell culture supernatants were screened based on the biochemical and biological properties of the rat plasma-derived TPO. As the result, TPO activities almost equal to the rat plasma-derived TPO were found in the culture supernatant of rat hepatocyte-derived cell lines McA-RH8994, HTC and H4-II-E, as well as in the culture supernatant of rat primary hepatocytes <Example

On the other hand, the expression vector pEF18S was constructed from 2 expression vectors pME18S and pEFBOS < Example 5>. Construction of this vector has rendered possible easy cloning of cDNA using a high efficiency expression vector having multiple cloning sites which can be used for the integration of inserts.

In addition to the above expression vector, pUC and pBR based plasmid vectors and λ based phage vectors are mainly used for the construction of cDNA libraries.

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The McA-RH8994 cells were cultured, homogenized CsCl density gradient centrifugation to obtain total RNA by adding guanidine thiocyanate solution and then subjected to <Example 6> Preparation of RNA can also be made by a hot phenol method, an acid guanidium phenol chloroform method and the like.

using oligo dT-immobilized latex particles, first strand cDNA After purifying poly (A)+ RNA from the total RNA was synthesized by the action of reverse transcriptase using recognition sequence has been added, followed by the synthesis expression vector pEF18S constructed in Example 5, which has been digested with Notl and EcoRI and then thus ligated oligo dT as a primer to which a restriction enzyme Notl of 2nd strand cDNA using RNase H and E.coli DNA polymerase I. An EcoRI adaptor was added to the thus obtained double-strand DNA, the resulting cDNA was ligated with a mammalian cell DNA was transformed into competent cells of an E. coli strain OH5 to construct a cDNA library < Example 7>.

Preparation (cloning) of rat TPO cDNA fragment by PCR O

amino acid sequence of the rat TPO which has been purified rom rat blood plasma, in order to synthesize degenerate Primers to be used may also be obtained based on an amino A DNA sequence was deduced from the partial acid sequence other than the position of the primers used herein. Highly degenerate primers may also be used without employing inosine. In addition, primers with reduced degeneracy can be designed making use of a codon having high primers to be used in the polymerase chain reaction (PCR), usage in rat (Wada et al., Nucleic Acids Res., vol.18, p.2367-2411, 1990).

When plasmid DNA was extracted from entire portion of the cDNA library prepared above and PCR was

carried out using the extracted DNA as a template, a band of about 330 bp was detected which was subsequently determined by nucleotide sequence analysis as a DNA fragment (A1 ragment) that encodes a portion of the rat TPO <Example 8>.

Screening of rat TPO cDNA by PCR, sequence of rat TPO cDNA and confirmation of TPO activity 6

out using each pool of plasmid DNA as a template and primers fragment, bands considered to be specific were detected in 3 The cDNA library prepared above was divided into pools, each containing about 10,000 clones, and plasmid DNA was extracted from each of 100 pools. When PCR was carried containing about 900 clones, and plasmid DNA was purified from 100 sub-pools to carry out PCR in the same manner. As these pools was divided into sub-pools, each containing 40 clones, and finally each clone was screened by PCR in the same manner. As the result, a clone pEF18S-A2 α which seemed to the result, a specific band was detected in 3 sub-pools. One of newly synthesized based on the nucleptide sequence of the A1 pools. One of these 3 pools was divided into sub-pools, each encode the rat TPO cDNA was isolated <Examples 9 and 10>.

When nucleotide sequence of this clone was analyzed, it was revealed that it encodes the partial amino acid sequence of the protein purified from rat blood plasma, hus confirming a strong possibility that this clone contains the rat TPO cDNA <Example 10>.

clone pEF18S-A2a contains a cDNA which encodes the rat TPO PO activity was found in the culture supernatant of the As the result, it was confirmed that the When plasmid DNA was purified from the clone pEF18S-A2α obtained above and transfected into COS 1 cells, transfected cells. Example 11>.

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TPO mRNA expression in rat tissues was analyzed by PCR, and specific expression was detected in the brain, the liver, the small intestines and the kidney <Example 12>.

(F) Construction of human cDNA library

Based on the results of Examples 4 and 12, the liver was selected as the starting tissue for the cloning of human TPO cDNA. In consequence, a cDNA library was constructed using a commercial mRNA derived from normal human liver. Using pEF-18S as a vector like the case of the rat library, cDNA was synthesized by the same procedure as the case of rat to obtain a directionally cloned cDNA library by making use of restriction enzymes Nofl and EcoRI. The library prepared by introducing vector-ligated cDNA into E. coli DH5 contained about 1,200,000 clones <Example 13>.

(G) Preparation (cloning) of human TPO cDNA fragment by PCR

Several primers for PCR were synthesized based on the nucleotide sequence of the clone pEF18S-A2α which encodes the rat TPO cDNA. When cDNA was synthesized using a commercial mRNA derived from normal human liver and PCR was carried out using these primers and cDNA as a template, a band of around 620 bp was observed. When the nucleotide sequence was analyzed, it was revealed that this clone contains a DNA fragment which has a homology of about 86% with the rat TPO cDNA, thus confirming a strong possibility that this is a part of a gene which encodes human TPO <Example 14s.

(H) Screening of human TPO cDNA by PCR, sequence of human TPO cDNA and confirmation of TPO activity

pools and subjected to PCR, possible bands were detected in 3 purified from each of the colonies was subjected to PCR to amplified, divided into pools, each containing about 100,000 on the nucleotide sequence of the human TPO fragment obtained in Example 14, possible bands were detected in 3 clones, and plasmid DNA was purified from each of 90 subpools, each containing 30 clones, and plasmid DNA was purified from each of 90 sub-pools and subjected to PCR, possible bands were detected in 3 sub-pools. Thereafter, 90 colonies were isolated from one of these sub-pools and plasmid DNA clones, and plasmid DNA was extracted from each of 90 pools. When PCR was carried out using each pool of plasmid DNA notecules as a template and primers newly synthesized based pools. One of these pools was divided into sub-pools, each containing 5,000 clones, and plasmid DNA was purified from each of 90 sub-pools. When PCR was carried out using each pool of plasmid DNA as a template in the same manner, possible bands were detected in 5 sub-pools. When one of these pools was divided into sub-pools, each containing 250 sub-pools. When one of these pools was divided into sub-The human cDNA library prepared above was finally obtain a clone named HL34 < Example 15>.

When nucleotide sequence of plasmid DNA in this clone was analyzed, it was revealed that this clone contains a cDNA which has about 84% homology with the rat TPO cDNA nucleotide sequence <Example 16>.

When the cloned plasmid DNA was purified and transfected into COS 1 cells, TPO activity was found in the culture supernatant of the transfected cells. As the result, it was confirmed that this plasmid clone contains the cDNA which encodes the human TPO <Example 17>.

However, this cDNA seemed to be an artificial product of the cloning, because no termination codon was found in this clone and a poly A tail-like sequence was found on its

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which encodes the amino acid sequence excluding a corresponding portion to the poly A tail-like sequence. When the thus constructed vector was expressed in COS 1 cells, TPO In consequence, an expression vector was constructed activity was found in the resulting culture supernatant <Example 18>.

DNA fragment in the 3'-end region of the human TPO was obtained by PCR so as to analyze the structure of a fullength cDNA. The determination of the nucleotide sequence of his fragment revealed that it partly overlaps with cDNA carried by the clone HL34 obtained in Example 15. It was also expected that a full length human TPO cDNA may contain an open reading frame thereof and code for a protein consisting of 353 amino acids. Thus, it was suggested that the human TPO consisted of 353 amino acids including a signal sequence of 21 amino acids <Example 19>.

In addition to the above cloning method, cloning of plaque hybridization using a rat TPO cDNA fragment as a probe, making use of a library constructed using a pUC or pBR based plasmid vector, a \(\eta \) phage based vector or the like. In designing a degenerate probe; inosine may be used to decrease the degree of degeneracy. When an assay method which can detect TPO activity in a specific manner or with a high sensitivity is human TPO cDNA can be attained by colony hybridization or available, it is possible to carry out expression cloning making use of an expression library as used in the present invention.

However, since the content of human TPO-encoding RNA seems to be extremely low in the normal human liver as from the result of this Example was in a ratio of one per three million), the hybridization screening in which a synthesized bligonucleotide or a rat or a human TPO cDNA fragment is used be extremely difficult to effect, because the will be described later in Example 15 (the content calculated

number of clones or plaques to be treated becomes bulky and sensitivity and specificity of the hybridization method are inferior to those of the PCR method. As a matter of fact, the nventors of the present invention have carried out colony hybridization of two million clones in a normal human liver cDNA library prepared in Example 13 using a rat TPO cDNA fragment as a probe, but a human TPO cDNA clone was not able to be obtained.

Reconstruction of human normal liver-derived cDNA

Since the clone HL34 obtained in Example 15 seemed to contain an incomplete cDNA, a cDNA library was reconstructed using a commercial normal human liver-derived poly (A)+ RNA preparation to obtain a complete human TPO cDNA. The library (hTPO-F1) prepared by introducing vector-ligated cDNA into E. coli DH5 contained 1.0 x 10⁶ transformants <Example 20>. Screening of TPO cDNA clone, sequence determination and expression of human TPO cDNA, and confirmation of TPO 3

Primers for PCR were synthesized based on the in Example 14 and the nucleotide sequence (SEQ ID NO: 196) of nucleotide sequence (SEQ ID NO: 3) of a partial cDNA obtained a complete cDNA of human TPO predicted in Example 19.

constructed in Example 20 was divided into 3 pools (pool # 1 -3) . PCR was carried out using the plasmid DNA prepared from each pool as a template and the synthesized primers. As the The # 3 pool was then divided into subpools, each containing 15,000 transformants, and screening using PCR was carried out as above. As the result, amplification of DNA having the The human liver cDNA library (hTPO-F1) result, a DNA fragment having the expected size was amplified when the plasmid DNA prepared from the pool # 3 was used.

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colonies grew on each LB plate, and a replica plate was prepared from each of the thus inoculated plates. When PCR was carried out in the same manner as described above, expected size was found in 6 of the 90 pools. When one of these positive pools was divided into subpools, each containing 1,000 clones, and plasmid DNA was extracted and PCR was carried out in the same manner, DNA amplification was not observed. It was considered to be caused by a low recovery of he plasmid DNA due to poorer growth of the clone of interest han the other clones. Therefore, the original # 3 pool was spread on 100 LB plates in such an inoculum size that 4,100 using these DNAs extracted from colonies grown on the plate amplification of a band expected was observed in one of 100 subpools.

Two replica filters were prepared from the plate of his subpool, and colony hybridization was carried out using he labeled probe(the EcoRI/BamHI fragment of plasmid pEF18S-HL34). As the result, a single signal considered to be positive was observed. The colonies were collected from the original plate and inoculated again onto an LB plate. DNA samples were prepared from a total of 50 colonies grown on he plate and were subjected to PCR to finally obtain a clone were mainly used and combination of these methods increased or it. When the nucleotide sequence of the thus obtained clone had an open reading frame and the amino acid sequence of a completely coincided with the deduced amino acid sequence SEQ ID NO: 6) of human TPO. The nucleotide sequence was different from the deduced one (SEQ ID NO: 196) at 3 positions, named pHTF1 <Example 21>. In screening of cDNA clone, nethods such as hybridization, PCR, and expression cloning efficiency and sensitivity of the screening, or decreased labor protein considered to be encoded by the open reading frame pHTF1 was determined, it was revealed that the clone pHTF1 out these differences did not cause amino acid exchange. It

residues having a 21 amino acid signal sequence. When the confirmed that human TPO protein comprises 353 amino acid and transfected into COS1 cells, TPO activity was found in the plasmid DNA was prepared from the thus obtained clone pHTF1 culture supernatant <Example 23>.

plaque hybridization, sequence determination and Screening of human TPO chromosomal DNA by means of expression of 3

human TPO chromosomal DNA and confirmation of TPO activity

onto 18 NZYM plates in such an inoculum size that one plate noculated again onto NZYM plates in such an inoculum size A genomic library given from Prof. T. Yamamoto at contained 30,000 phage particles, and two replica filters were prepared from each of the thus prepared 18 plates. A human TPO DNA fragment (base position numbers 178 to 1,025 in SEQ ID NO: 7) was amplified by PCR and purified. Using the purified fragment [abeled with $^{32}\mathrm{P}$ as a probe, plaque hybridization was carried out. As the result, 13 positive signals were obtained. Plaques were collected from the original plates and that 1,000 plaques were formed on each plate. Two replica ilters were prepared from each of the resulting plates to carry out plaque hybridization under the same conditions he Gene Research Center, Tohoku University was inoculated described above. As the result, positive signals were detected on all filters of the 13 groups. A single plaque was recovered DNA samples thus prepared from the 13 clones were checked or the presence of the cDNA coding region by PCR. Five of the predicted from the cDNA.In consequence, one of these clones clone 1HGT1) was selected and analyzed by Southern biotting rom each of the resulting plates to prepare phage DNA. Phage 13 clones seemed to contain entire amino acid coding region using the aforementioned probe). Since a single band of about - 46 -

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the clone JHGT1 was digested with HindIII and subjected to agarose gel electrophoresis. The 10 kb band was excised from the gel, purified, and subcloned into a cloning vector pUC13 to 10 kb was observed in the case of the HindIII digestion, DNA of finally obtain a clone named pHGT1 < Example 24>.

clone pHGT1 was determined, chromosomal DNA carried by the clone was found to contain an entire coding region of the the coding region coincided completely with that of the When the nucleotide sequence of the thus obtained protein predicted in Example 19, and nucleotide sequence of predicted nucleotide sequence (SEQ ID NO: 196).

acid-encoding exon contained 4 introns and the nucleotide sequence was different from that of the complete cDNA In addition, a region corresponding to the amino carried by clone pHTF1 obtained in Example 21 (SEQ ID NO: 7).

When nucleotide sequences of 4 remaining clones among the 5 chromosomal DNA clones independently selected by the screening were determined, it was found that 2 of the 4 clones were identical to the clone pHGT1 and other 2 clones were the same as the clone pHGT1 except that they had a different nucleotide within 3' noncoding region as observed vith the clone pHTF1 <Example 25>.

An EcoRI fragment in the thus obtained plasmid clone pHGT1 was ligated with an expression vector pEF18S to obtain human TPO expression plasmid pEFHGTE. When the plasmid DNA (pEFHGTE) was prepared and transfected into COS1 cells, TPO activity was found in the culture supernatant <Example 26>

expression in COS1 cells and confirmation of TPO Preparation of human TPO deletion derivatives, their activity 3

he plasmid clone pHT1-231 obtained in Example 18 as a numan TPO protein could exhibit its biological activity even after the removal of its carboxyl-terminal third. Thus, in order to further analyze the biologically active portions, deletion derivative experiments were performed. A series of expression plasmids were prepared by PCR using the DNA of The results of Example 18 and 22 revealed that emplate and synthesized oligonucleotides as primers.

Expression plasmids which contained DNA coding erminal region of the TPO protein, that is, deletion 163 amino acid sequence, were obtained. When the plasmid DNA was prepared from each of clones and transfected into COS1 cells, TPO activity was detected in each of the culture or human TPO deletion derivatives lacking the carboxylderivatives coding for positions 1-211, 1-191, 1-171 and 1supernatant < Example 27>.

derivatives having respective deletion of N-terminal side designed and the activity after their expression in COS 1 cells when N-terminal side amino acid residues were deleted up to he 7 position or C-terminal side amino acid residues were When derivatives having a series of deletion of Cterminal amino acid residues up to the 151 position and other amino acid residues up to the 6, 7 and 12 position were was measured in order to analyze the TPO activity-bearing egion further in detail, the activity became undetectable deleted up to the 151 position. <Examples 28 and 29>

Derivatives of a protein having its TPO biological insertion or addition) cDNA coding the protein. Methods such as PCR, site directed mutagenesis and chemical synthesis can activity can be obtained by modifying (deletion, substitution, be used for the modification.

Expression of human TPO cDNA in CHO cell and purification of TPO. E

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An expression vector, pHTP1, was constructed which can express cDNA coding for a deduced amino acid sequence of human TPO as shown in SEQ ID NO: 6, in animal cells. <Example 30>

The TPO cDNA region in pHTP1 was used to construct a vector pDEF202-hTPO-P1 for its expression in CHO cell. <Example 31>

As a result of the transfection of the vector into CHO cells and subsequent selection of transfected cells, the transfected cells in which the expression vector carrying human TPO cDNA has been integrated into chromosomes of CHO cells were obtained. <Example 32>

In Example 32, large scale cultivation was carried out for a human TPO producing CHO cell line (CHO 28-30 cell, resistant to 25 nM MTX) which has been prepared by transfecting the human TPO expression plasmid pDEF202-hTPO-P1 into CHO cells. <Example 55>

From the culture supernatant 100 L, human TPO was purified. < Example 56>

Alternatively, human TPO was purified from the culture supernatant in Example 55 in a different manner. < Example 57>

(N) Expression of human TPO cDNA in X63.6.5.3. cells and confirmation of the activity

Using the TPO cDNA region encoded in the plasmid pBLTEN prepared in Example 30, an expression vector BMCGSneo-hTPO-P1 was constructed for use in X63.6.5.3. cells. Example 33

When X63.6.5.3. cells were transfected with this vector, a transformant cell was obtained in which the human cDNA-encoding expression vector was integrated into its

chromosome. When this cell was cultured, TPO activity was detected in the culture supernatant. <Example 34>

(O) Large quantity expression of human TPO in COS 1 cells, and purification, molecular weight measurement and biological characteristics thereof

The expression vector pHTP1 prepared in Example 30 was transfected into COS 1 cells to obtain a large quantity (a total of about 40 liters) of culture supernatant containing the expressed product. < Example 35>.

Purification of TPO was carried out from about 7 liters of the COS 1 cell serum-free culture supernatant prepared in Example 35 containing the expression vector pHTP1-derived TPO. It was able to obtain high activity TPO through the steps of a hydrophobic interaction chromatography, a cation exchange column chromatography, a WGA column chromatography and a reverse phase column chromatography. eExample 36>

The TPO thus partially purified from the COS 1 cell culture supernatant was subjected to molecular weight measurement and biological characteristics analysis.

Examples 37 and 38>

(P) Expression of human TPO in E. coli

A vector pGEX-2T/hT(1-174) was constructed for use in the expression of a fusion protein (to be referred to as "GST-TPO(1-174)") of glutathione-S-transferase and human TPO (amino acid residues of positions 1 to 174) in *E. coli.* In this case, a portion of the human TPO cDNA nucleotide sequence (about half the area of the 5'-side) was exchanged with *E. coli* preference codons. <Example 39>

GST-TPO(1-174) was expressed in *E. coli*, the esulting cells were disintegrated and then the GST-TPO(1-174) contained in the precipitation fraction was solubilized.

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Also, a vector pCFM536/h6T(1-163) was constructed for use in the expression in *E. coli* of a mutation type human TPO protein (to be referred to as "h6T(1-163") in which the 1-position Ser residue and the 3-position Ala residue of human TPO (amino acid residues of positions 1 to 163) were respectively replaced by an Ala residue and a Val residue and, at the same time, a Lys residue and a Met residue were respectively added to the -1 and -2 positions. Entire portion of the human TPO cDNA nucleotide sequence (corresponding to 1 to 163 amino acid residues) carried by this vector was exchanged with *E. coli* preference codons. <Example

The h6T(1-163) was expressed in *E. coli*, the resulting cells were lysed and then solubilization of h6T(1-163) contained in the precipitated fraction and conditions for the refolding of the protein were examined, thereby succeeding in partially purifying a protein containing a TPO amino acid sequence as designed. It was confirmed that the protein shows TPO activity in the rat CFU-MK assay system. <Examples 43 and 44>

In addition, a vector pCFM536/hMKT(1-163) was constructed for use in the expression in *E. coli* of a mutation type human TPO protein (to be referred to as "hMKT(1-163)") in which a Lys residue and a Met residue were respectively added to the -1 and -2 positions. The hMKT(1-163) was expressed in

E. coli in the same manner as described in Example 43, and the expression protein was subjected to SDS-PAGE, transferred on a PVDF membrane and then subjected to N-terminal amino acid sequence analysis, thereby confirming that this protein contains the designed amino acid sequence. <Example 52>

In addition, the vector pCFM536/hMKT(1-332) for expression in *E. coli* of a mutation type human TPO protein in which a Lys residue is added at the position-1 and Met residue at the position-2 of human TPO (amino acid positions 1 to 332) (referred to as "hMKT (1-332)") respectively was constructed. The hMKT (1-332) was expressed within *E. coli* in the same manner as in Example 42, and its expression was confirmed by Western blotting using the anti-human TPO peptide antibody prepared in Example 45 set forth below. <Example 66>

(Q) Preparation of anti-TPO peptide antibody and anti-TPO peptide antibody column

Rabbit polydonal anti-TPO peptide antibodies were prepared by synthesized peptides corresponding to three partial regions of the rat TPO amino acid sequence determined in Example 10. It was confirmed that these antibodies can recognize rat and human TPO molecules. Furthermore, peptides corresponding to 6 partial regions in the amino acid sequence of human TPO shown in SEQ ID NO: 6 (or SEQ ID NO: 7) were synthesized and then employed to prepare rabbit polyclonal anti TPO peptide antibodies. The resultant antibodies were confirmed to recognize human TPO. <Example 45>

It is possible to purify TPO by an affinity column chromatography using a column packed with certain molecules having binding affinity for TPO, such as anti-TPO antibodies, TPO receptors and the like, are bound. In consequence, an anti-TPO antibody column was firstly prepared by binding the anti-TPO peptide antibody obtained in Example 45. <Example 46>

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and molecular weight and biological characteristics making use of anti-TPO peptide antibody column, thereof

Using a culture supernatant from the transfection of COS 1 cells with the expression vector pHTP1 as a starting naterial, partially purified TPO was obtained and applied to the anti-TPO antibody column. Since the TPO activity was found in the adsorbed fraction, this fraction was further subjected to a reverse phase column chromatography to purify the protein and examine its molecular weight and biological characteristics. <Example 47>

Confirmation of the activity of partially purified sample of human TPO expressed in COS 1 cells <u>(S</u>

namely the TPO sample purified up to the step of Capcell Pak Cl 300A column from a culture supernatant obtained by transfecting COS 1 cells with the expression vector pHTP1, was checked for its biological activities to determine if it The TPO-active fraction purified in Example 36, exerts a platelet number increasing action in the living body. <Example 48>

exchange column from 33 liters of a culture supernatant prepared by transfecting COS 1 cells with the expression vector pHTP1 was checked for its biological activities to find Also, a crude TPO fraction obtained by a cation that it increases platelet number in the body. <Example 49>

Expression of human TPO chromosomal DNA in CHO cells and confirmation of the activity E

A vector pDEF202-ghTPO was constructed for use in the expression of human TPO chromosome in CHO cells. <Example 50>

When this vector was introduced into CHO cells, a transformant cell was obtained in which the human TPO chromosomal DNA-encoding expression vector was integrated into its chromosome. When this cell clone was cultured, TPO activity was detected in the culture supernatant. <Example

ö Partial purification and activity confirmation mutation type human TPO expressed in E. coli 3

The mutation type human TPO derived from the pCFM536/h6T(1-163), and expressed in E. coli was subjected to refolding using guanidine hydrochloride and glutathione to ind that the thus obtained h6T(1-163) exerts a platelet <Example 53> nucleotide sequence-encoding number increasing action in the living body. human TPO

The mutation type human TPO derived from the that the thus purified h6T(1-163) increases platelet number in pCFM536/h6T(1-163), and expressed in E. coli was subjected to refolding using sodium N-lauroyl sarcosinate and copper sulfate and then to a cation exchange chromatography to find clone human TPO nucleotide sequence-encoding the living body. <Example 54>

In addition, the refolding and purification of the mutation type human TPO, h6T (1-163) was conducted using another procedures. <Examples 60 and 61>

Expression of human TPO cDNA in insect cells and identification of TPO activity 3

A recombinant virus for the expression of human TPO in insect cells was prepared <Example 58> and expressed - 54 -

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in the insect cell Sf 21, and subsequently the TPO activity was identified in the culture supernatant. <Example 59>

(W). Expression of human TPO (amino acid positions 1 to 163) in CHO cells and purification thereof

The vector pDEF202-hTPO163 for expression in CHO cells of the human TPO protein (referred to as "hTPO163") having amino acids 1 to 163 in the human TPO amino acid sequence shown in SEQ ID NO: 7 was constructed. The expression vector pDEF202-hTPO163 was transfected into CHO cells and thus-obtained hTPO163-producing CHO cell line was cultured in a large scale. From the supernatant of the culture hTPO163 was purified. <Examples 62 to 65>

(X) Preparation of substitution derivatives of human TPO

Arg-25 and Glu-231 of human TPO have been substituted by Asn and Lys, respectively, was prepared as well as a derivative (referred to as "09/TPO") wherein His-33 has been replaced by Thr.

A plasmid coding for each derivative was transfected into COS7 cells which were then cultured. TPO activity was detected in the supernatant of the culture. <Example 67>

Derivatives wherein one amino acid of h6T(1-163) has been substituted by an other amino acid were prepared by using E. coli expression system. TPO activity was found in each derivative. < Example 94>

 (Y) Preparation of insertion or deletion derivatives of human IPQ

An amino acid is inserted into or deleted from hTPO163 to prepare its insertion or deletion derivatives, and thereafter COS7 cells are transfected with plasmids encoding the resultant derivatives. In the supernatant of the COS7 cell cultures, the TPO activity was detected. <Example 68>

A method for the measurement of TPO activity (an in vitro assay system) used in the present invention is described in the following as "Reference Example".

(Z) Preparation and purification of anti-human TPO antibodies

Polyclonal antibodies are prepared using peptide fragments of human TPO < Example 69 and subsequently utilized in Western blot analysis < Example 70 and in the construction of anti-TPO antibody affinity columns < Example 71 sequences.

(AA) In vivo activity of human TPO

Purified human TPO is administered to mice with induced thrombocytopenia and changes in platelet counts are monitored versus a control group. <Example 72>, through <Example 79>. Pharmaceutical compositions are described as potential useful in treatment of thrombocytopenia. <Example 80> through <Example 89>.

BB) IPO activity as a function of MLP receptor binding

An assay is provided wherein TPO activity is defined in terms of specific binding interaction with the Mpl receptor. <Example 90> through <Example 93>.

Reference Example

 A. Rat megakaryocyte progenitor cell assay (rat CFU-MK, assay) system (liquid culture system) 26

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1977). This phenomenon can be observed already in small and mononuclear acetylcholinesterase-positive cells which are considered to be positioned between CFU-MK and recognizable in response to the increment of the size of megakaryocytes 1981). In addition, it is known that such phenomenon is Megakaryocytes incorporate and store serotonin in p.672, 1984). The amount of serotonin incorporated increases specific for only megakaryocytic cells among bone marrow CFU-MK cells (Gplib/Illa+ CFU-MK fraction which will be described later) are cultured in the presence of samples to be dependent process (Fedorko, Lab. Invest., vol.36, pp.310 - 320, megakaryocytes (Bricker and Zuckerman, Exp. Hematol., vol.12, Schick and Weinstein, J. Lab. Clin. Med., vol.98, pp.607 - 615, cells (Schick and Weinstein, J. Lab. Clin. Med., vol.98, pp.607 -615, 1981). In the present assay system, highly enriched rat assayed, and incorporation of 14C-serotonin (14C-hydroxy ryptamine creatine sulphate; 14C-5HT) into megakaryocytes cytoplasmic dense granules through an active, energygrown from CFU-MK is measured.

Advantages of this assay system are that indirect influences of contaminated cells (for example, formation of Meg-CSF activity by contaminated cells stimulated by a certain substance other than the factor of interest, or formation of a certain factor by contaminated cells, which undergoes a combined action with the factor of interest) can be reduced, because the percentage of CFU-MK in cells of GpIIb/IIIa+ CFU-MK fraction is markedly high (see the following [Assay method]) while the number of contaminating cells is small. In addition, appropriate culture conditions can be maintained for a relatively long period, because the total number of cells seeded per one well is small. Another advantage is that a number of large-size mature megakaryocytes grown from CFU-MK in the presence of an

active sample during a culture period can visually detected under a phase contrast microscope, thus rendering possible qualitative judgement of the presence and degree of the activity. Results of quantitative judgement correspond well to the results of quantitative determination based on the incorporation of ¹⁴C-serotonin. Therefore, reliability of the quantitative determination can be further improved by joint use of the qualitative judgement.

ssav method:

Highly enriched rat CFU-MK cells (Gpllb/Illa⁺ CFU-MK fraction) used in the assay were prepared by slightly modifying the method of Miyazaki *et al.* (*Exp. Hematol.*, vol.20, pp.855 - 861, 1992).

the megakaryocyte separation medium reported by Levine and suspension. The thus prepared bone marrow cell suspension is density; 1.050 g/ml/1.063 g/ml/1.082 g/ml) which is Pharmacia) with HATCH solution, and centrifuged at 20°c at 100 x g for 20 minutes. After the centrifugation, cells between the 1.063 g/ml and 1.082 g/ml density layers are ecovered. After washing, the recovered cells are suspended in Briefly, femur and tibia are removed from Wister suspension as described. A suspending medium (a solution (Path-O-Cyte 4; Seikagaku Kogyo Co., Ltd.) and Ca2+ and Mg2+ree Hanks' balanced salt solution: to be referred to as "HATCH 1977) is used for the preparation of the bone marrow cell ayered over a Percoll discontinuous density gradient solution prepared by diluting Percoll solution (manufactured by ats (male, 8 to 12 weeks of age) to prepare bone marrow cell consisting of 13.6 mM trisodium citrate, 11.1 mM glucose, 1 mM adenqsine, 1 mM theophylline, 10 mM HEPES (pH 7.25), 0.5% bovine serum albumin (to be referred to as "BSA" hereinafter) solution" hereinafter) which is a slightly modified version of Fedoroko (Levine and Fedoroko, Blood, vol.50, pp.713 - 725,

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scove's modified Dulbecco culture medium (to be referred to as "IMDM culture medium" hereinafter) containing 10% fetal calf serum (to be referred to as "FCS" hereinafter), put in 100mm tissue culture plastic dishes and cultured at 37.C for 1 cells are recovered and cultured again in plastic dishes at 37.C for 1 hour. Non adherent cells are suspended in HATCH

hour in a 5% CO2 incubator. After incubation, nonadherent

petri dishes to which a mouse monoclonal anti-rat platelet

Spllb/Illa antibody, P55 antibody (Miyazaki et al., Thromb. Res.,

solution and incubated for 1 hour on 100-mm bacteriological

After incubation, nonadherent cells are removed by thorough washing with HATCH solution, and the remaining cells adsorbed to the immobilized P55 antibody are detached by pipetting and collected. In general, 3-4 x 10^5 cells are

vol.59, pp.941 - 953, 1990) has previously been adsorbed

CFU-MK fraction" hereinafter), and it is known that this cell raction contains about 5 to 10% of CFU-MK based on the measurement by a colony assay system in the presence of a saturated concentration of rat IL-3. A hybridoma capable of producing the aforementioned P55 antibody has been deposited under the deposit No. FERM BP-4563 in National Institute of Sioscience and Human Technology, Agency of Industrial Science and Technology, Ministry of International Trade and Industry on

obtained from one rat. The thus obtained cell fraction contains

righly enriched rat CFU-MK (to be referred to as "GpIlb/IIIa+

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scintillation counter. otu

In this instance, almost the same results as those of the 14C-serotonin incorporation assay can be obtained when acetylcholinesterase activity in the aforementioned cell ysate is measured according to the procedure of Ishibashi and Burstein (Blood, vol.67, pp.1512 - 1514, 1986)

Standard samples:

Firstly, blood plasma of thrombocytopenic rats was prepared in the following manner, for use in the preparation of standard samples.

> raction are suspended in IMDM culture medium containing 10% FCS and dispensed in portions of 104 cells per well of a 96 well tissue culture plate. Each well is further supplied with a standard sample (will be described later in detail), or a sample to be assayed, thereby adjusting the final medium volume to 200 µl/well. The thus prepared plate is put in a CO2 incubator

and incubated for 4 days at 37 .C. On 4th days of incubation,

Next, cells of the thus obtained GpIIb/IIIa+ CFU-MK

-ebruary 14, 1994.

Normal male Wistar rats (7 to 8 weeks old) were rendered thrombocytopenic by intravenous administration of the aforementioned P55 antibody with a dose of 0.5 mg per animal, twice with an interval of about 24 hours. About 24 hours after the second administration, blood was collected rom the abdominal aorta under ether anesthesia, using a 10 ml

once again. A 200 µl portion of 2% Triton X-100 is added to shaken on a plate mixer for approximately 5 to 10 minutes to yse the cells thoroughly. A 150 µl portion of the thus hours before the completion of culturing, and the final ncubation is continued at 37.C. After 3-hr incubation, the plate is centrifuged at 1,000 rpm for 3 minutes, and the esulting supernatant fluid is removed by suction. A 200 µl and the plate is centrifuged to wash the plate by removing the esulting supernatant fluid. This washing step is repeated the thus obtained pellet of cells in each well, and the plate is obtained cell lysate is transferred into a commercial capshaped solid scintillator (Ready Cap; manufactured by Beckman) which is subsequently left overnight at 50°C in an oven to dry the lysate. On the next day, the Ready Cap is put a glass vial to measure radioactivity of ¹⁴C by a liquid portion of PBS containing 0.05% EDTA is added to each well, $0.1~\mu Ci~(3.7~KBq)$ of ^{14}C -serotonin is added to each well



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capacity syringe in which 1 ml of 3.8% (v/v) trisodium citrate solution has been included as an anti-coagulation agent. The thus collected blood sample was dispensed into plastic centrifugation tubes and centrifuged at 1,200 x g for 10 minutes to recover blood plasma fraction. The thus recovered blood plasma fraction was again centrifuged at 1,200 x g for 10 minutes, and the resulting blood plasma fraction was ecovered, taking care to avoid the pellet consisting of cells, platelets and the like, and pooled. (The blood plasma obtained reated TRP (standard sample C), WGA-Agarose column active raction (standard sample W) or Phenyl Sepharose 6 FF/LS column active fraction (standard sample P) obtained from TRP in accordance with the procedure of Example 1 were dialyzed extensively against a sufficient volume of IMDM culture in such a manner is referred to as "TRP" hereinafter.) nedium and used as assay standards.

In the rat TPO purification process described in Example 1, the standard sample C was used in the early stage, but was changed to the standard sample W halfway thereafter and then to the standard sample P. Specific activity of the standard sample C was defined tentatively as 1, and relative activities of the standard samples W and P were calculated based on the definition. Relative activity of each sample to be assayed was determined by comparing the dose-response curve as n where its activity is n times higher than that of the of the standard sample to those of the samples to be assayed, and relative activity of the sample to be assayed was defined standard sample C.

B. Colony assay system

In this assay system, bone marrow cells are cultured in a semi-solid culture medium in the presence of a sample to be assayed, and the Meg-CSF activity is measured by

counting the number of megakaryocyte colonies formed by the proliferation and differentiation of CFU-MK.

Assay method:

In the case of the use of unseparated rat bone marrow cells and the like (a)

procedure of Jackson (Blood, vol.42, pp.413 - 421, 1973). When manufactured by DIFCO) was put in a 35-mm tissue culture cultured at 37.C in a CO2 incubator. In general, the number of disks are detached from the dishes and placed on glass slides (76 mm x 52 mm). To dry each agar disk, a piece of 50-_m acetylcholinesterase staining solution prepared according the sufficient staining of megakaryocytes is confirmed, the agar slides are washed with water, dried, subjected to postwashed with water and then air-dried. Megakaryocyte colonies A 1 ml final volume of IMDM culture medium containing un-separated rat bone marrow cells, cells obtained in each separation step of GpIIb/IIIa+ CFU-MK of the aforementioned assay system A, or cells of Gpllb/Illa+ CFU-MK fraction, and 10% FCS, 2 mM glutamine, 1 mM sodium pyruvate, 50 µM 2-mercaptoethanol and 0.3% agar (AGAR NOBLE, plastic dish and, after solidification at room temperature, cells per one dish is adjusted to $2-4 \times 10^5$ in the case of unseparated bone marrow cells, 2-5 x 104 in the case of cells at the step of the Percoll density gradient or of the adherence Jepletion, or 0.5-2 x 103 in the case of cells of GpIIb/Illat CFU-MK fraction. After 6 to 7 days of the culture, the agar nylon mesh and filter paper are placed, in that order, over the gel. The thus dried agar slides are heat-fixed for 5 minutes at 50.C on a hot plate and soaked for 2 to 4 hours in an staining with Harris' hematoxylin solution for 30 seconds. tre defined as 3 or more tightly grouped acetylcholinesterasepositive cells.

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In the case of the use of unseparated mouse bone marrow e

The colony assay can be performed in the same manner as the above procedure (a) using 2-4 imes 10⁵ cells per one dish.

marrow cells or human cord blood cells In the case of the use of human bone <u>ပ</u>

Human bone marrow cells or human cord blood cells may be used as such, or as CFU-MK fractions enriched by the following manner.

over Lymphoprep (manufactured by Daiichi Kagaku Co., Ltd.) and centrifuged, and the resulting interface leucocyte fraction is From the thus recovered cell fraction, cells to which biotinylated monoclonal antibodies specific for human Colony assay of human cells case of the use of rat bone marrow cells, except that 3-5 x 103 cells of the CD34+DR+ CFU-MK fraction are used in one used instead of 10% FCS. The culturing period for the Firstly, bone marrow fluid or cord blood is layered surface antigens (CD2, CD11c and CD19) bind are removed with avidin-linked magnetic beads. The cells removed by the magnetic beads method are mainly B cells, T cells, macrophages and a part of granulocytes. The remaining cells are stained with FITC-labeled anti-CD34 antibody and PElabeled anti-HLA-DR antibody, and then a CD34- and HLA-DR-ELITE manufactured by COULTER). CFU-Mit cells are can be carried out in the same manner as the aforementioned dish and a mixture of 12.5% human AB plasma and 12.5% FCS is formation of megakaryocyte colonies is 12 to 14 days. For the positive fraction is recovered using a cell sorter (for example, concentrated in this fraction (to be referred to as "CD34+DR+ CFU-MK fraction" hereinafter). recovered.

detection of human megakaryocytes, immunological staining of anti-alkaline phosphatase antibody method with a mouse monoclonal antibody specific for a megakaryocyte surface antigen GpIIb/IIIa (for example, Teramura et al., Exp. Hematol., vol.16, pp.843 - 848, 1988), and colonies each consisting of 3 megakaryocytes is carried out by an alkaline phosphataseor more megakaryocytes are counted.

Assay system with a human megakaryoblast cell line (M-OZe assay C

proliferate in response to GM-CSF, IL-3, SCF, IL-2 and the like Since these cells also respond to TPO, they are applicable to a Avanzi et al., J. Cell. Physiol., vol.145, pp.458 - 464, 1990). M-07e cells, a human megakaryoblast cell line, substitutive assay system for the rat CFU-MK assay system.

Assay method:

of culturing. After completion of the culturing, the cells are thereby adjusting the final volume to 200 µl/well. The thus prepared plate is put in a 5% CO2 incubator and incubated for 3 thymidine is added to each well 4 hours before the completion collected on a glass fiber filter using a cell harvester to M-07e cells maintained in the presence of GM-CSF are recovered, washed thoroughly and then suspended in IMDM culture medium containing 10% FCS. The resulting M-07e cell suspension is dispensed in portions of 10^4 cells into wells of a 96 well tissue culture plate, and each well is further supplied with a standard sample or a sample to be assayed, days at 37°C. After 3 days of culture, 1 μCi (37 KBq) of $^3 \text{H-}$ measure 3H radioactivity with a liquid scintillation counter for example, Beta Plate manufactured by Pharmacia).

which a mouse pro-B cell line is used An assay system (Ba/F3 assay) in

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Mouse pro-B cell line Ba/F3 is known as a cell line which proliferates in response to IL3, IL4 and the like (Palacios et al., Cell, vol.41, pp.727 - 734). Since its substrain L3 and IL4 but also TPO, it can be applied to an assay system as a substitute for the rat CFU-MK assay or the human megakaryoblastic cell line-aided assay (M-07e assay) is as L-3 are recovered, washed three times with IMDM and plate of a density of 1 x 104 cells per well, each well is urther supplemented with a standard TPO solution or a sample The resulting plate is incubated for 2 to 3 days in a 5% CO2 or additional 4 hours. Finally, the cells are collected on a system, almost all of cells cultured in media lacking TPO However, cells cultured in media containing TPO activity show and incorporate 3H-thymidine. In addition, results of this BF-TE22 is capable of cell proliferation in response to not only ollows. Briefly, BF-TE22 cells growing in Iscove's modified DME medium(IMDM:GIBCO) in the presence of 1 ng/ml of mouse suspended in IMDM medium containing 10% FCS. Next, the cell suspension is dispensed into wells of a 96 well tissue culture thymidine is added to each well and the culturing is continued radioactivity with a liquid scintillation counter. In this assay activity die and therefore do not incorporate ³H-thymidine. active proliferation in a TPO concentration-dependent manner to be tested and adjusted to a final volume of 200 µl/well. On the 2nd or 3rd day, 1 µCi (37 KBq) of ³H· glass fiber filter using a cell harvester to measure ³H ncubator.

The following examples are provided to further illustrate he present invention.

assay are parallel to those of the M-07e and CFU-MK assays.

EXAMPLE 1-1

Purification of rat TPO from blood plasma of thrombocytopenic rats induced by anti-platelet antibody administration

Preparation of blood plasma of thrombocytopenic rats induced by anti-platelet antibody administration

about 1,000 rats in accordance with the procedure described in TRP was prepared as the purification source from the aforementioned <Reference Example> A, rat megakaryocyte precursor cell (CFU-MK) assay system.

Purification of rat TPO from TBP

about 1,000 rats on an assumption that TPO content in TRP would be one per three million at the most, and a little less Initially, purification was carried out using TRP of than 1 pmole of partially purified rat TPO was obtained. Though it is difficult in general, an attempt was made to analyze its partial amino acid sequences and three partial amino acid sequences were obtained but with a low accuracy. These sequences coincided with or are close to the amino acid sequence of a serine protease inhibitor (SPI) produced in the liver. It was unclear from these results whether TPO has a structure similar to that of the serine protease inhibitor or the sequences are derived from those of contaminated proteins other than TPO. Using these uncertain sequences, cloning of TPO gene from a rat cDNA library was carried, but potential Thereafter, ntensive studies were carried out based on an assumption that such a failure was due to low purity and insufficient amount of the sample analyzed. In order to obtain a purified final sample necessary for the amino acid analysis, purification was carried out in accordance with a process described in the ollowing <Example 1-2>. Surprisingly, it was estimated from results of <Example 1-2> that the TPO content in TRP or XRP will be described below) was such a extremely small amount as one per one hundred million to one billion of total plasma TPO-encoding genes could not be obtained.

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proteins, so that its complete purification was extremely difficult.

EXAMPLE 1-2

Puritication of rat TPO from blood plasma of thrombocytopenic rats induced by total body X-ray or γ -ray irradiation [Preparation of blood plasma of thrombocytopenic rats induced by total body X-ray or γ -ray irradiation]

Normal male Wister rats (7 to 8 weeks old) were rendered thrombocytopenic by whole body irradiation with X-rays or γ -rays at a sublethal dose (6 to 7 Gy). Blood was collected from the rats on the 14th day after irradiation and a blood plasma fraction (to be referred to as "XRP" hereinafter) was prepared in the same manner as the aforementioned procedure for the preparation of TRP.

In this case, XRP (ca. 8 L) prepared from a total of about 1,100 rats was used as the purification source.

Purification of rat TPO from XRP

The blood plasma sample of about 1,100 rats was too large to process at one time because its total protein content reached 493,000 mg. In consequence, the blood plasma sample was divided into 11 lots, each corresponding to about 100 rats, in the following purification steps (1) to (4). In the subsequent purification steps (5) to (7), the sample was divided into 6 batches. In and after the purification step (8), a sample crudely purified from the total blood plasma of about 1,100 rats was used. In the following, typical example of each purification step is described in the case of a lot (XW9) and a batch (XB6).

In all of the purification steps, TPO activity was measured using the rat CFU-MK assay system described in <Reference Example>. Unless otherwise noted, all purification

steps were carried out at 4_C, excluding the reverse phase chromatographies, and Superdex 75pg gel filtration in the presence of surfactant, which were carried out at room temperature. Protein assay was carried out by a Coomassie dye binding assay (a reagent system manufactured by PIERCE, catalog No. 23236X) or a method using bicinchoninic acid (a reagent system manufactured by PIERCE, catalog No. 23225).

An outline of the purification is shown in Table 1.

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Table 1 Purification of rat plasma TPO

538	2022000	8687	797	S-200HR F2 (high mol. TPO)	
811	1020000	20000	15	S-200HR F3 (low mol. TPO)	< Gel filtmlion>
142	2339000	∠ †8	79 <i>L</i> Z	Phenyl-Sepharose 6 PF/LS	< Hydrophobic>
443	3834000	506	4536	TSK-gel AF-Blue 650MH	< Triazine dye>
230	1987000	135	12030	DROIGSA-ADW	< Lectin>
513	000¥0L	6	314400	Q-Sepharose PP	< อฐิตรศิวxอ ติดโ >
001	009+98	Z	480300	Ca-Treated plasma/G25	• •
_		<u> </u>	493000	Total plasma	
(%)			(gm)		•
recovery	κιίνίις	activity	protein		
Activity	IsloT	Relative	IntoT	Step	

Low molecular TPO (from S-200HR F3)

				(under non-reduction cond.)	
67	220300	14900000	4100.0	12% 2D2-PAGE	< Electrophoresis >
77	009861	4890000	96£0.0	Capcell Pak CI	(Reverse phase>
SE	298400	800000	0.3730	YMC CN-AP	< Reverse phase>
43	371000	130000	2.8540	YMC Protein-RP prop	< Reverse phase>
911 (%)	0007001	20000	(3m) 50.3300	S-200HR F3 (low mol. TPO)	< noimatin lab>
recovery	activity	activity	nistorq		
Aictivity	IstoT	Relative	IsloT	Step	

High molecular TPO (from 5-200HR F2)

				non-reduction cond.)	nabau)
411	000066	330000000	0.0030	15% SDS-PAGE	< Electrophoresis >
971	1260000	20000002	0.0630	Capcell Pak Cl	< Reverse phase>
OS .	434000	00000L	0.6200	YMC CN-AP	< Reverse phase>
536	2041000	1750000	0991.1	Superdex 75pg/CHAPS	< Gel filtration>
300	2588000	227000	11.4000	mol. TPO) YMC Protein-RP prep	< Reverse phase>
733 (%)	2012000	7840	(gm) 257.0000	S-200HR F2 (high	<ดงเกิดเกิด (จอ)
recovery	Activity	activity	nistorq		
Activity	Lotal	Relative	Total	Step	

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protease inhibitor treatment of rat blood plasma Calcium chloride treatment, centrifugation and Ξ

In the case of lot No. XW9

Nalgene). Calcium chloride powder was added to each tube to a total protein, 40,686 mg) corresponding to about 100 rats, which had been stored at -80°C, was thawed and dispensed 4.C, the resulting mixture was centrifuged at 8,000 rpm for 50 minutes. To the resulting TPO-active supernatant fluid No. 010-10393) to a final concentration of 1 mM. The thus XRP (742 ml; protein concentration, 54.8 mg/ml; nto polypropylene centrifugation tubes (manufactured by inal concentration of 100 mM. After overnight incubation at (742 ml; protein concentration, 54.9 mg/ml; total protein, 40,740 mg) was added a protease inhibitor p-APMSF ((paminodiphenyl)methanesulfonyl fluoride hydrochloride, manufactured by Wako Pure Chemical Industries, Ltd., catalog obtained sample was used in the following Sephadex G-25 column buffer exchange step.

In this manner, calcium chloride/p-APMSF treatment of the whole XRP derived from about 1,100 rats (total volume, 8,184 ml; total protein, 493,000 mg) was to about 100 rats, and carrying out treatment of these lots one effected by dividing it into 11 lots, each lot being equivalent by one. The resulting 11 samples were processed separately in the subsequent Sephadex G-25 column step.

(2) Sephadex G-25
 se In the case of lot No. XW9

treatment of the step (1) (742 ml; protein concentration, 54.9 mg/ml; total protein, 40,740 mg) was applied at a flow rate of 40 to 70 ml/min to a Sephadex G-25 column (manufactured by The supernatant fluid obtained after the calcium

Pharmacia Biotech, catalog No. 17-0033-03; diameter, 11.3 elution of protein was discarded. When UV absorption was active protein fraction exchanged by 20 mM Tris-HCI, pH 8 cm; bed height, 47 cm) which had been equilibrated in advance with 20 mM Tris-HCl, pH 8. Elution was carried out with the same buffer. A 1,300 ml portion of the eluate before the detected in the eluates, fractions were pooled until electric conductivity reached 500 µS/cm, thereby recovering a TPO-(1,377 ml; protein concentration, 27.56 mg/ml; total protein, 37,882 mg; protein yield, 93%). Relative activity of TPO in the raction was found to be 2.3.

As the result of the Sephadex G-25 column reatment of the whole lots of samples, a total of 21,117 ml of the Sephadex G-25 column TPO-active fraction was obtained (total protein, 480,300 mg; average relative activity, 2; total activity, 864,600).

(3) Q-Sepharose FF <strong anion exchange chromatography> In the case of lot No. XW9

by the Sepharose G-25 treatment (1,377 ml; protein concentration, 27.5 mg/ml; total protein, 37,841 mg; relative by Pharmacia Biotech, catalog No. 17-0510-01; diameter, 5 with 20 mM Tris-HCI (pH 8) and a fraction F1 which passed ml; protein concentration, 0.98 mg/ml; total protein, 3,870 mg; relative The TPO-active fraction obtained in the above step activity, 2.3) was applied to a Q-Sepharose FF (manufactured cm; bed height, 27 cm) at a flow rate of 40 ml/min, eluted through the column was pooled (3,949 activity, 0). <u>(Z</u>

Next, the buffer was changed to 20 mM Tris-HCI (pH 8) containing 175 mM NaCl to elute a TPO-active fraction F2 (4,375 ml; protein concentration, 5.36 mg/ml).

ml; protein concentration, 3.9 mg/ml; total protein, 4,783 mg; relative Finally, a fraction F3 (1,221

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activity, 3.8) was eluted with 20 mM Tris-HCI (pH 8) containing 1,000 mM NaCl. Total protein in the TPO-active fraction F2 was found to be 23,440 mg, and protein yield of F2 by this step was 61.9%. In addition, relative activity of TPO was increased to 6.8. As the result of the application of the whole lots FF, a total of 35,842 ml of the Q-Sepharose FF TPO-active of the Sephadex G-25 TPO-active fractions to the Q-Sepharose fraction F2 was obtained (total protein, 314,384 mg; average relative activity, 9; total activity, 2,704,000).

Wheat germ agglutinin (WGA)-Agarose <le>slectin_affinity_chromatography> **4**

In the case of lot No. XW9

The TPO-active fraction F2 obtained in the above step (3) by the Q-Sepharose FF treatment was divided into three portions and applied to a WGA-Agarose (manufactured by which passed through the column was pooled (9,336 ml; Honen Corp., catalog No. 800273; diameter, 5 cm; bed height, 22.5 cm) at a flow rate of 5 ml/min, and eluted with Dulbecco's isotonic phosphate buffer (DPBS). A fraction F1 protein concentration, 2.30 mg/ml; total protein, 21,407 mg; relative activity, 6.9).

Next, the buffer was changed to 20 mM sodium resulting eluates were pooled and concentrated using an phosphate buffer (pH 7.2) containing 0.2 M N-acetyl-Dglucosamine (GlcNAc, manufactured by Nacalai tesque, catalog No. 005-20), 150 mM NaCl and 0.02% sodium azide, and the ultrafiltration unit (Omega Ultrasette, nominal molecular weight cutoff of 8,000; manufactured by Filtron), thereby obtaining a WGA-Agarose adsorbing TPO-active fraction F2 (2,993 ml; protein concentration, 0.376 mg/ml).

Total protein in the TPO-active fraction F2 was found to be 1,125 mg, and protein yield of F2 by this step was

2 In addition, relative activity of TPO was increased 101. The thus obtained F2 fraction was stored at -80°C.

Agarose, a total of 33,094 ml of the WGA-Agarose TPO-active As the result of the application of the whole lots of the Q-Sepharose FF TPO-active F2 fractions to the WGAfraction F2 was obtained (total protein, 15,030 mg; average relative activity, 132; total activity, 1,987,000).

TSK-gel AF-BLUE 650 MH <triazine dye affinity chromatography>

in the case of batch No. XB6

starting from 215 rats-equivalent XRP were combined as a The WGA-Agarose adsorbing TPO-active fraction of lot No. XW8 and the WGA-Agarose adsorbing TPO-active fraction F2 of lot No. XW9 obtained in the above step (4) batch No. XB6 (5,947 ml, protein concentration, 0.388 mg/ml; total protein, 2,319 mg; relative activity, 150).

to make a 6,132 ml of solution having a final NaCl concentration of 0.822 M, and the resulting solution was applied at a flow rate of 7 ml/min to a TSK-gel AF-BLUE 650 A 5,974 ml portion of the combined sample was MH column (TOSOH CORP., catalog No. 08705; diameter, 5 cm; bed height, 23 cm) which had been equilibrated in advance with mixed with 0.85 moles of NaCl for 1,000 ml (296.76 g in total) 20 mM sodium phosphate buffer (pH 7.2) containing 1 M NaCl.

After completion of the application, protein was eluted using 20 mM sodium phosphate buffer (pH 7.2) eluates were pooled and concentrated using an ultrafiltration unit (Omega Ultrasette, nominal molecular weight cutoff of 8,000), thereby obtaining a passed-through fraction F1 (543 ml; protein concentration, 2.05 mg/ml; total protein, 1,112 containing 1 M NaCl at a flow rate of 10 ml/min. The resulting

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active fraction F2 (1,427 ml; protein concentration, 0.447 Next, the elution buffer was changed to 2 M NaSCN to obtain an eluted TSK-gel AF-BLUE 650 MH adsorbing TPO-

Total protein in the TPO-active fraction F2 was found to be 638 mg, and protein yield of F2 by this step was In addition, relative activity of TPO was increased to 27.5%. of all of the fractions to the TSK-gel AF-BLUE 650 MH, a total of 10,655 ml of the TSK-gel AF-BLUE 650 MH TPO-active fraction F2 was batches of the WGA-Agarose adsorbing TPO-active F2 obtained (total protein, 4,236 mg; average relative activity, As the result of the application 905; total activity, 3,834,000).

Phenyl Sepharose 6 FF/LS < hydrophobic interaction chromatography> 9

In the case of batch No. XB6

ng/ml; total protein, 638 mg; relative activity, 1,500) was mixed with 1.5 moles of ammonium sulfate powder per 1,000 ml (282.2 g in total) to make a 1,581 ml of solution having a A 1,424 ml portion of the TSK-gel AF-BLUE 650 MH TPO-active fraction F2 (1,424 ml; protein concentration, 0.447 final ammonium sulfate concentration of 1.35 M.

containing 0.8 M ammonium sulfate at a flow rate of 10 The resulting solution was applied at a flow rate of 7 ml/min to a Phenyl Sepharose 6 FF (Low Sub) column (Pharmacia Biotech, catalog No. 17-0965-05; diameter, 5 cm; bed height, 10 cm) which has been equilibrated in advance with 50 mM sodium phosphate buffer (pH 7.2) containing 1.5 M elution was carried out using 36 mM sodium phosphate buffer ml/min The resulting eluates (about 3,160 ml) were pooled and concentrated using an ultrafiltration unit (Omega ammonium sulfate. After completion of the application,

Ultrasette, nominal molecular weight cutoff of 8,000), thereby obtaining a fraction F1 (485 ml; protein concentration, 0.194 mg/ml; total protein, 94.2 mg; relative activity, 0).

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active fraction F2 (about 3,500 ml). The eluted fraction was nominal molecular weight cutoff of 8,000) and a sample for assay was taken out. At this stage, protein concentration and total protein of the TPO-active fraction F2 (220 ml) were found to be 1.45 mg/ml and 319 mg, respectively, and protein yield of F2 by this step was 50.0%. Relative activity of TPO sodium phosphate buffer (pH 7.2) to obtain an eluted TPOconcentrated using an ultrafiltration unit (Omega Ultrasette, Next, the elution buffer was changed to 20 was found to be 1,230.

As the result of the application of all of the fractions to the Phenyl Sepharose 6 FF/LS, a total of 1,966 ml of the Phenyl Sepharose 6 FF/LS TPO-active fraction F2 was batches of the TSK-gel AF-BLUE 650 MH TPO-active F2 obtained (total protein, 2,762 mg; average relative activity, 847; total activity, 2,339,000).

Sephacryl S-200 HR <pel filtration chromatography> In the case of batch No. XB6 (cf. Fig. 1) <u>S</u>

315 mg; relative activity, 1,230) was mixed with 144.8 ml of 5 M NaCl solution to make a 362 ml solution having a final NaCl concentration of 2 M, and the resulting solution was concentrated to about 50 ml using an ultrafiltration unit with YM 3 membrane (76 mm in diameter, manufactured by Amicon The Phenyl Sepharose 6 FF/LS TPO-active fraction F2 (217 ml; protein concentration, 1.45 mg/ml; total protein,

To this was added the same volume (50 ml) of 8 M M NaCl and 4 M urea as final concentrations. This was concentrated to about 80 ml, made into a sample of 88.78 ml urea, thereby obtaining about 100 ml of a solution containing 1

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and then applied to a Sephacryl S-200 HR column (manufactured by Pharmacia Biotech, catalog No. 17-0584-01; 7.5 cm in diameter and 100 cm in bed height).

Thereafter, elution was effected with DPBS at a llow rate of 3 ml/min, and the eluates after a void volume of 1,200 ml were collected in 45 ml portions in 60 polypropylene were stored at -85°C until the Sephacryl S-200 HR treatment of all samples derived from 1,100 rats was completed. Based lubes. Assay was carried out every two tubes and the rest on the assay results, fractions of the batch No. XB6 were grouped as follows.

(F1) tube numbers 1 to 15 (fractions around void volume;

molecular weight, 94,000 or more)

(F2) tube numbers 16 to 26 (molecular weight, 94,000 to

33,000)

(molecular weight, 33,000 to (F3) tube numbers 27 to 44

3.000)

(molecular weight, 3,000 or less) (F4) tube numbers 45 to 55

ractions obtained by Phenyl Sepharose 6 FF/LS were stored samples were thawed and concentrated using an nanufactured by Amicon Corp.) to obtain the following two samples. The concentrated sample of the TPO-active fraction In this manner, all batches of the TPO-active F2 separately subjected to Sephacryl S-200 HR, assayed and stored at -85 C. After completion of the S-200 HR treatment of all batches and just before the subsequent operation of everse phase chromatography (YMC-Pack PROTEIN-RP), these ultrafiltration unit with YM 3 membrane (76 mm in diameter, F2 obtained by the Sephacryl S-200 HR is referred to as "high molecular TPO sample F2" hereinafter, and that of F3 as "tow nolecular TPO sample F3".

of convenience, and the term "high molecular" and "low molecular TPO sample F3 are pooled fractions of different The high molecular TPO sample F2 and the low elution areas in the gel filtration chromatography as a matter molecular" therefore may not mean their true molecular

Molecular weight	High molecular TPO sample F2 94,000 - 33,000	Low molecular TPO sample F3 33,000 - 3,000
lotal volume Total volume	3,420 mi 293 ml	6,480 ml 280 ml
(after concentration)	262 ma	51.3 ma
Average relative activity 7,838	rity 7,838	20,000
Total activity	2,055,000	1,020,000

molecular TPO sample F2 are separately subjected to the The low molecular TPO sample F3 and the high subsequent purification steps.

Purification of the low molecular TPO sample F3 is described in the following steps (8) to (11).

YMC-Pack PROTEIN-RP <reverse phase chromatography> <u>@</u>

espectively. Insoluble materials formed during the The low molecular TPO sample F3 (total protein, obtained in the above step (7) was mixed with a solvent A containing 0.025% TFA) to obtain a solution having a total volume of 508.63 ml and final propanol, TFA and protein concentrations of about 20%, 0.012% and 0.0989 mg/ml, and the resulting supernatant was divided into two 254.3 ml 20,000; total activity, 1,007,000; total volume, 274 ml) (0.025% trifluoroacetic acid (TFA)) and a solvent B (1-propanol preparation of this solution were separated by centrifugation, 50.3 mg; protein concentration, 0.184 mg/ml; relative activity,

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25.2 mg protein) portions and applied at a flow rate of 2 (manufactured by YMC, catalog No. A-PRRP-33-03-15; 3 cm in n advance with 30% B. The precipitate resulting from the centrifugation was dissolved in 20 ml of 20 mM sodium acetate (pH 5.5) containing 5 mM CHAPS (3-[3ml/min to a column packed with YMC-Pack PROTEIN-RP diameter and 7.5 cm in bed height) which has been equilibrated

cholamidopropyl)dimethylammonio]-1-propane suffonate;

manufactured by Dojindo Laboratories, catalog No. 75612-03-

3) and also applied to the column.

(solvents A:B = 3:1) was passed through the column to obtain a After the sample loading, about 50 ml of a solvent and the eluates were collected in 36 polypropylene tubes in 10 ml portions. This process was repeated again for the remaining sample using the same collection tubes, thus passed-through fraction. Next, a developing program (120 minutes of linear gradient from 30% B to 45% B) was started, obtaining a total of 36 fraction tubes each containing 20 ml eluate. The passed-through fraction was directly concentrated to 20 ml using an ultrafiltration unit with YM 3 membrane (76 nm in diameter, manufactured by Amicon Corp.).

raction and the 20 ml fractions of tube numbers 1 to 36 was mixed with 20 µl of 5% BSA, dried by evaporation, dissolved in was found in the fractions of tube numbers 17 to 27 (36.0 to A 0.1 ml portion of each of the passed-through 0.25 ml of IMDM assay culture medium and then assayed to specify TPO-active fractions. As the result, the TPO activity 43.0% propanol concentration range) which were combined and used as a low molecular TPO sample F3-derived YMC-Pack at -85°C until its use in the subsequent YMC-Pack CN-AP step. PROTEIN-RP TPO-active fraction F2.

ow molecular TPO sample F3-derived YMC-Pack PROTEIN-RP

active fraction F2

0.0130 mg/ml 2.85 mg 130,000 371,000 220 ml Protein concentration Relative Activity **Fotal** activity Total protein Total volume

YMC-Pack CN-AP < reverse phase chromatography>

obtained in the above step (8) was mixed with 0.6 ml of 50% glycerol and concentrated to 1.8 ml. The concentrate was inally adjusted to a volume of 5 ml containing 20% or less of sample F3-derived YMC-Pack PROTEIN-RP TPO-active fraction F2 (total protein, 2.79 mg; protein concentration, 0.0130 mg/ml; relative activity, 130,000; total activity, 36,300) A 214.9 ml portion of the low molecular TPO propanol and about 6% of glycerol.

513; 6 mm in diameter and 250 mm in bed height) which has from 15% B to 25% B, and 65 minutes of linear gradient from in the column. Since the same polypropylene tubes were used in the following column operation (0.555 mg protein and 1 ml volume for each operation). Each of the thus divided samples was applied at a flow rate of 0.6 ml/min to a column packed been equilibrated in advance with 15% B, using 0.1% TFA as the B. After the application, propanol concentration was increased 5th) operation, 1 ml of a solution having the same composition excluding protein was applied to the column and developed in the same manner to recover TPO activity retained o pool the eluates of the 6 operations, a total of 44 tubes The concentrate was divided into 5 portions for use with YMC-Pack CN-AP (manufactured by YMC, catalog No. APsolvent A and 0.05% TFA-containing 1-propanol as the solvent 25% B to 50% B was carried out. After completion of the final



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μl of 5% BSA, dried by evaporation, dissolved in 0.24 ml of active fractions. As the result, strong TPO activity was found A 30 µl portion of each of the thus obtained ractions (1/240 portion of each fraction) was mixed with 20 IMDM assay culture medium and then assayed to specify TPOin the fractions of tube numbers 28 to 33 (37.0 to 42.0% propanol concentration range) which were combined and used as a low molecular TPO sample F3-derived YMC-Pack CN-AP main TPO-active fraction FA.

ow molecular TPO sample F3-derived YMC-Pack CN-AP TPOactive

fraction FA

43.20 ml	0.00863 mg/ml	0.373 mg	800,000	298,400
Total volume	Protein concentration	Total protein	Relative Activity	Total activity

stinal reverse phase 300 (10) Capcell Pak C1 chromatography>

A 43.12 ml portion of the 43.20 ml low molecular 0.372 mg; protein concentration, 0.00863 mg/ml; relative activity, 800,000; total activity, 297,500) obtained in the above step (9) was mixed with 0.2 ml of 50% glycerol and TPO sample F3-derived TPO-active fraction FA (total protein, concentrated to obtain 0.1 ml of glycerol solution. This solution was mixed with 2 ml of a solution of solvent A (0.1% TFA): solvent B (1-propanol containing 0.05% TFA) = 85:15 (15% B) to prepare 2.1 ml of sample containing about 14% propanol, about 4.8% glycerol and 0.177 mg/ml of protein. The thus prepared sample was applied to a column packed with Capcell Pak C1 300A (manufactured by Shiseido Co., Ltd., catalog No. C1 TYPE:SG300A; 4.6 mm in diameter and

250 mm in bed height) which has been equilibrated in advance with 15% B and developed by 65 minutes of linear gradient from 27% B to 38% B at a flow rate of 0.4 ml/min. The eluates were collected in 72 polypropylene tubes in 0.6 ml portions.

A 3 µl portion of each of the thus obtained ractions (1/200 portion of each fraction) was mixed with 20 μl of 5% BSA, the medium was exchanged to 225 μl of IMDM assay culture medium and the thus 75 times diluted fraction was assayed.

and treated at 95°c for 5 minutes with 10 µl of a nonreducing sample buffer for SDS-PAGE. The thus treated sample was subjected to SDS-PAGE using a 15-25% SDS-polyacrylamide and then stained with a silver staining kit of 2D-Silver Stain-II Low Molecular Weight Markers (manufactured by Daiichi Pure Chemicals Co., Ltd., catalog No. 181061; to be referred to as A 1 µl portion of each fraction (1/600 fraction) was aliquotted for use in electrophoresis, dried by evaporation precast gel (manufactured by Dailchi Pure Chemicals Co., Ltd.) 'DAIICHI" (manufactured by Dailchi Pure Chemicals Co., Ltd., catalog No. 167997; to be referred to as "silver staining kit" nereinafter). As to molecular weight markers, [DAIICHI]-III 'DPCIII" hereinafter) were used.

As the result of the above assay, significant TPO activity was found in the fractions of tube numbers 35 to 43 ractions from tube numbers 36 to 42 (30.5 to 32.0% propanol 30.0 to 32.5% propanol concentration range). Of these, concentration range) were combined and used as the main TPOactive fraction FA. The results are shown in Fig. 2.

1,890,000 and a total activity of 193,600. Examination of the SDS-PAGE patterns of the TPO-active fraction numbers 36 to When the protein content deduced from the chromatogram was compared with the assay results, the thus obtained fraction was found to have a total protein of 39.6 µg, a protein concentration of 9.4 µg/ml, a relative activity of

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12 revealed the presence of a band whose staining density In addition, apparent molecular weight of this non-reduced band was found to be 17,000 to 19,000 when compared with those of the standard molecular weight proteins on the same gel, which were reduced, thus revealing that this band is a strong candidate of correlates with the activity strength.

from electrophoresis gel <15% SDS-PAGE> (11) Extraction of TPO-active protein

Example of the analysis of TPO-active fraction FA

Of the 4,200 µl of the low molecular TPO sample (1/1680 fraction) for use in silver staining were transferred 37.c for 1 hour and then allowed to stand still for 18 hours at F3-derived TPO-active fraction FA (total protein, 39.6 µg; protein concentration, 9.4 µg/ml; relative activity, 4,890,000; iotal activity, 193,600) obtained in the above step (10), 5.5 µl (1/764 fraction) for use in active protein extraction and 2.5 µl into respective sample tubes, dried by evaporation, reacted with 10 µ of a nonreducing sample buffer for SDS-PAGE at room temperature, thereby effecting the SDS reaction.

out at 4 c in accordance with a usually used procedure to silver staining were immediately cut out with a knife, put into a fixing solution and then stained using the Pre-stained Low Range Marker (manufactured by Bio-Rad Laboratories, Inc., 161-0305) and the aforementioned DPCIII were used as molecular weight markers. Making use of a microslab gel, 15% SDS-PAGE of these samples was carried (Laemmli, Nature, vol.227, pp.680 - 685, 1970). Atter completion of the electrophoresis, gel portions to be subjected aforementioned silver staining kit.

Separately from this, all molecular weight range of gel to be used for the detection of the activity was cut, the

with a knife into 34 slices, and each gel slice having a width of 1.5 to 2.5 mm was smashed by a slightly modified method of Each gel slice thus smashed into minute fragments was mixed with 0.3 ml of an extraction buffer (20 mM Tris-HCl, pH 8, 500 mM NaCl, 0.05% BSA) and shaken for 6 hours at 4 c to effect Kobayashi (Kobayashi, M., Seikagaku (Biochemistry), vol.59, no.9, 1987, published by the Japanese Biochemical Society). extraction.

at 4 . c, the resulting mixture was transferred into an Ultra 5.8) to a final concentration of 20 mM. After 1 hour of shaking Free C3GV 0.22 µm filtration unit (manufactured by Millipore for 15 minutes to remove precipitated SDS and recover the resulting supernatant fluid. The filtrate was transferred into an Ultra Free C3-LGC ultrafiltration unit (nominal molecular weight cutoff of 10,000, manufactured by Millipore Corp., UFC3 -GC 00) and centrifuged at 3,000 x g (7,000 rpm). When To this was added 500 mM potassium phosphate (pH Corp., UFC3 OGV 0S) and centrifuged at 1,000 \times g (4,000 rpm) volume of the concentrated solution reached about 50 µl, 300 ul of 20 mM sodium phosphate buffer (pH 7.2) was added and again subjected to the ultrafiltration.

hereafter, a similar step was repeated to exchange to the which was subsequently sterilized and subjected to TPO The ultrafiltration using 300 µl of 20 mM sodium phosphate buffer was repeated twice to remove remaining SDS. assay culture medium for preparation of a sample (300 µl) activity measurement.

Three protein bands were detected clearly by the silver staining which showed apparent molecular weights of about 17,000 to 19,000, 14,000 and 11,000 based on the DPCIII molecular weight markers. In the previous step (10), a band having a correlation between activity strength and staining density and naving an apparent molecular weight of about 17,000 to

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19,000 was observed in the electrophoresis of the Capcell Pak C1 column TPO-active fraction. In the experiment of this step (11), TPO activity was found also in the band having an apparent molecular weight of about 17,000 to 19,000 (cf. Fig.

of Capcell Pak C1 300A column to a detectable level on the electrophoresis gel. Based on the silver-stained density of the band obtained by 15% SDS-PAGE, the amount of the candidate IPO protein (an apparent molecular weight of about 17,000 to 19,000) in the total TPO-active fraction was determined to be that the TPO-active protein was purified in the active fraction On the basis of the above results, it was confirmed about 1.7 µg.

Purification of the high molecular TPO sample F2 described in the following steps (12) to (15).

(12) YMC-Pack PROTEIN-RP < reverse phase chromatography>

The high molecular TPO sample F2 (total protein, obtained in the above step (7) was mixed with 95.8 ml (1/3 volume of the sample) of a solvent B (1-propanol containing 3.025% TFA) to obtain a solution having a total volume of 383 ml and final propanol, TFA and protein concentrations of about 0.025% TFA was used for a solvent A. Insoluble materials ormed during the preparation of input sample were separated by centrifugation, and the resulting supernatant fluid was at a flow rate of 2 ml/min to a column packed with YMC-Pack 15; 3 cm in diameter and 7.5 cm in bed height) which has been equilibrated in advance with 30% B. The precipitate resulting from the centrifugation was dissolved in 10 ml of 20 mM 7,840; total activity, 2,015,000; total volume, 287 ml) 25%, 0.006% and 0.671 mg/ml, respectively. A solution of divided into six 62.3 ml (42.8 mg protein) portions and applied 257 mg; protein concentration, 0.894 mg/ml; relative activity PROTEIN-RP (manufactured by YMC, catalog No. A-PRRP-33-03-

sodium acetate (pH 5.5) containing 5 mM CHAPS and also applied to the column.

solvents A:B = 3:1) was passed through the column to obtain a passed-through fraction. Next, a developing program (120 and the eluates were collected in 24 polypropylene tubes in 15 ml portions. This process was repeated for the 6 divided through fraction and the tube number 1 fraction were combined and directly concentrated to 90 ml using an ultrafiltration unit minutes of linear gradient from 30% B to 45% B) was started, samples using the same collection tubes, thus obtaining a total with YM3 membrane (76 mm in diameter, manufactured by After the sample toading, about 50 ml of a solvent of 24 fraction tubes each containing 90 ml eluate. The passed-Amicon Corp.).

raction plus tube number 1 fraction and the fractions of tube he result, the TPO activity was found in the fractions of tube which were combined and used as a high molecular TPO sample A 0.3 ml portion of each of the passed-through evaporation, dissolved in 0.30 ml of IMDM assay culture medium and then assayed to specify TPO-active fractions. As numbers 10 to 15 (34.0 to 39.5% propanol concentration range) This fraction was stored at -85.c until its use in the numbers 2 to 24 was mixed with 10 µl of 5% BSA, dried by F2-derived YMC-Pack PROTEIN-RP TPO-active fraction F2. subsequent YMC-Pack CN-AP step. High molecular TPO sample F2-derived YMC-Pack PROTEIN-RP

active fraction F2

540 ml	n 0.021 mg/ml	11.4 mg	227,000	2.588.000
Total volume	Protein concentration	Total protein	Relative Activity	Total activity

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chromatography in the presence of CHAPS> (13) Superdex 75 pg <gel filtration

and concentrated by evaporation. To this was added 18 ml of incubation at 4.c. Thereafter, the first sample was applied to (manufactured by Pharmacia Biotech, catalog No. 17-1070-01; ml portion of each sample (protein concentration, 0.468 A 538.2 ml portion of the high molecular TPO 20 mM CHAPS, followed by stirring and a subsequent 41 hour a column packed with HiLoad 26/60 Superdex 75 pg 2.6 cm in diameter and 60 cm in bed height) and eluted with DPBS containing 5 mM CHAPS at a flow rate of 1 ml/min. A 4 sample F2-derived YMC-Pack PROTEIN-RP TPO-active fraction relative activity, 227,000; total activity, 2,565,000) obtained n the above step (12) was mixed with 0.6 ml of 50% glycerol F2 (total protein, 11.3 mg; protein concentration, 0.021 mg/ml mg/ml; protein, 1.86 mg) was applied to the column.

operation were collected in 5 ml portions in 45 tubes, thus inally obtaining 45 fractions each containing 30 ml eluate repeated 6 times by dividing the entire YMC-Pack PROTEIN-RP TPO-active fraction into 6 portions. The eluates of each In this case, the Superdex 75 column operation was after completion of the 6 column operations.

3,000) which were combined and used as a high molecular TPO evaporation, dissolved in 0.25 ml of IMDM assay culture medium and then assayed to specify TPO-active fractions. As the result, the TPO activity was found in the fractions of tube numbers 13 to 31 (molecular weight range of from 78,000 to A 0.1 ml portion of each of the thus obtained fractions was mixed with 10 µl of 5% BSA, dried by sample F2-derived Superdex 75 pg TPO-active fraction F2.

High molecular TPO sample F2-derived Superdex 75 pg TPOactive

fraction F2

0.00216 mg/ml 1,750,000 2,041,000 1.17 mg 540 ml Protein concentration Relative Activity **Fotal** activity Total volume Total protein

(14) YMC-Pack CN-AP <reverse phase chromatography>

sample F2-derived Superdex 75 pg TPO-active fraction F2 540 1,750,000; total activity, 1,943,000) obtained in the above propanol containing 0.05% TFA) and applied at a flow rate of 3.6 ml/min to a column packed with YMC-Pack CN-AP and 250 mm in bed height) which has been equilibrated in advance with 15% B, using a solvent A (0.1% TFA) and the solvent B. After the application, propanol concentration was A 513.2 ml portion of the high molecular TPO ml (molecular weight, 78,000 - 3,000) (total protein, 1.11 mg; protein concentration, 0.00216 mg/ml; relative activity, step (13) was mixed with 1/10 volume of a solvent B (1manufactured by YMC, catalog No. AP-513; 6 mm in diameter increased from 15% B to 25% B, and 65 minutes of linear gradient from 25% B to 50% B was carried out.

epeated three times. Since the same 44 polypropylene tubes Prior to the commencement of the YMC-Pack CN-AP operation twice. In other words, the column operation was column chromatography, a 1/20 portion of the total input sample was subjected to a trial operation in order to confirm proper recovery of the activity. Thereafter, the remaining 19/20 portion was divided into 2 samples to carry out the vere used to pool the eluates of the 3 operations, a total of 44 ubes each containing 3.6 ml eluate were obtained

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High molecular TPO sample F2-derived YMC-Pack CN-AP TPO active

fraction FA

25.20 ml	0.0246 mg/ml	0.620 mg	700,000	434,000
Total volume	Protein concentration	Total protein	Relative Activity	Total activity

<final reverse phase 300A chromatography> Pak (15) Capcell_

FA (total protein, 0.606 mg; protein concentration, 0.0246 mg/ml; relative activity, 700,000; total activity, 424,000) A 24.66 ml portion of the 25.20 ml high molecular TPO sample F2-derived YMC-Pack CN-AP TPO-active fraction obtained in the above step (14) was mixed with 0.4 ml of 50% glycerol and concentrated by evaporation.

In this way, 2 ml of a concentrated sample was Capcell Pak C1 300A (manufactured by Shiseido Co., Ltd., catalog No. C1 TYPE:SG300A; 4.6 mm in diameter and 250 mm in bed height) which has been equilibrated in advance with 15% B, using a solvent A (0.1% TFA) and a solvent B (1-propanol 0.303 mg/ml. This was applied to a column packed with obtained with a propanol concentration of a few percent, a glycerol concentration of 10% and a protein concentration of

containing 0.05% TFA), and developed by 65 minutes of linear The eluates were collected in 72 polypropylene tubes in 0.6 µl gradient from 27% B to 38% B at a flow rate of 0.4 ml/min. portions.

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A 0.75 µl portion of each of the thus obtained ractions (1/800 portion of each fraction) was mixed with 20 ul of 5% BSA, the medium was exchanged to 225 µl of IMDM assay culture medium and the thus 300 times diluted fraction was assayed. A 2 µl portion of each fraction (1/300 fraction) was sampled for use in electrophoresis, dried by evaporation and reated at 95 c for 5 minutes with 10 µl of a nonreducing sample buffer for SDS-PAGE. The thus treated sample was subjected to SDS-PAGE using a 15-25% SDS-polyacrylamide precast gel (manufactured by Daiichi Pure Chemicals Co., Ltd.) and then stained using the aforementioned silver staining kit. DPCIII described in the foregoing was used as molecular weight markers.

activity was found in the fractions of tube numbers 33 to 39 As the result of the above assay, significant TPO (29.5 to 31.5% propanol concentration range). Of these, ractions of tube numbers 34 to 39 (30.0 to 31.5% propanol active fraction FA. Examination of the SDS-PAGE pattern of he main TPO-active fraction revealed the presence of a band within an apparent molecular weight range of 17,000 to 19,000 under nonreducing condition, which was similar to the ase of the low molecular TPO sample F3-derived TPO-active concentration range) were combined and used as the main TPOwhose staining density correlates with the activity strength, raction described in the aforementioned step (10).

Analysis of partial amino acid sequences of purified rat TPO

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Amino acid sequence analysis of the rat TPO candidate protein in the Capcell Pak C1 300A column TPOactive fraction FA obtained in the purification step (10) of Example 1 was carried out in accordance with the procedure of wamatsu (Iwamatsu et al., "Shin Kiso Seikagaku Jikken-ho New Basic Biochemical Experiments)*, vol.4, pp.33 - 84, pub. Maruzen; Iwamatsu, A, Selkagaku (Biochemistry), vol.63, no.2, pp.139 - 143, 1991; Iwamatsu, A., Electrophoresis, vol.13, pp.142 - 147, 1992). That is, the sample was subjected to SDS-PAGE and transferred electrically onto a polyvinylidene difluoride (PVDF) membrane. After reduction and S-alkylation stepwise restrictive enzymatic hydrolysis in situ with three proteases, and the resulting peptide fragments on each digestion step were separated and purified by reverse phase of the protein on the PVDF membrane, the thus treated protein was digested into peptide fragments by systematic and chromatography and analyzed for their amino acid sequences by a high sensitivity amino acid sequence determination method. The following describes this process in detail.

Example of the analysis of the TPO candidate protein in the low molecular TPO sample F3-derived Capcell Pak C1 300A **IPO fraction FA**

Concentration of Capcell Pak C1 300A column TPO fraction FA (tube numbers 36 to 42)

F3-derived Capcell Pak C1 300A column TPO-active fraction activity, 193,600), 4,151 µl (98.8% of the total fraction) was used in the amino acid sequence analysis. The total protein FA (tube numbers 36 to 42) obtained in the purification step (10) of Example 1 (total protein, 39.6 µg; protein concentration, 9.4 µg/ml; relative activity, 4,890,000; total Of the 4,200 µl of the low molecular TPO sample deduced from the chromatogram was 39.1 µg, of which about

SDS-PAGE and having an apparent molecular weight of about 1.6 µg was the TPO candidate protein stained with silver after 17,000 to 19,000.

This sample was mixed with glycerol and concentrated by evaporation to obtain 5 µl of a glycerol obtaining about 25 µl of sample containing 200 mM Tris-HCl (pH 8.0), 50 mM Tris-HCI (pH 6.8), 1.1% SDS, 2 mM EDTA, 0.02% solution. To this were added a non-reducing buffer for SDS-PAGE and 1 M Tris-HCl (pH 8) in order to adjust its pH, thereby promophenol blue and 30% glycerol.

emperature for 14 hours and then treated at 60°c for 5 The thus prepared sample was maintained at room minutes, in order to effect proper SDS reaction without excess eating.

Electrophoresis (2)

Micro-slab gels (4.0% acrylamide stacking gel and 15% acrytamide separation gel) were prepared and SDS-PAGE was carried out at room temperature for 2 hours at a constant current of 12.5 mA and then at 17.5 mA. Pre-stained Low Range Marker (Bio-Rad, 161-0305) and DOCIII were used as Immediately after the electrophoresis, the resulting protein molecules were ransferred on a PVDF membrane (see the following step). molecular weight markers.

A portion of the sample was also used in another electrophoresis using 15-25% polyacrylamide precast gel Multi Gel 15/25 manufactured by Dalichi Pure Chemicals Co., Ltd., catalog No. 211072) under non-reducing conditions or after reducing the sample with dithiothreitol (DTT). When the esulting gel was stained using the silver staining kit described in the foregoing, it was confirmed that molecular weight of the protein band expected to be TPO was about 19,000 under a reducing condition and purity of the TPO andidate protein of Capcell Pak C1 300A column was a few

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percent. In addition, since mobility of the band varied under non-reducing and reducing conditions, it was suggested that the candidate protein contains at least one disulfide bond.

Transfer of protein on PVDF membrane by electroblotting and detection of band 3

Protein transfer on a PVDF membrane (ProBlott, manufactured by Applied Biosystems; catalog No. 400994) was carried out for 1 hour at a constant current of 160 mA (11 to 17 V) using a semi-dry transfer apparatus (Model KS-8460, manufactured by Marysol). A solution consisting of 0.3 M Tris and 20% methanol (pH 10.4) was used as the anolyte, a solution consisting of 25 mM Tris and 20% methanol (pH 10.4) as the transfer membrane solution and a solution consisting of 25 mM fris, 40 mM aminocaproic acid and 20% methanol (pH 10.4) as the cathlyte.

When the thus transferred membrane was stained with a Ponceau S staining solution (0.1 g of Ponceau S and 1 ml of acetic acid in 100 ml water), a plurality of bands were detected and one of these bands was confirmed to be that of the candidate protein having a molecular weight of about 19,000. This band was cut out for use in the subsequent peptide fragment formation steps.

Peptide fragment formation, peptide mapping and aming acid sequence analysis €

alkylated after reduction on the PVDF membrane, stepwise In order to perform systematic fragmentation of the TPO candidate protein which has been transferred and Srestrictive enzymatic hydrolysis was carried out using the following three proteases.

irst digestion: Lysyl endopeptidase (Achromobacter lyticus m497-1, manufactured by Wako Pure Chemical Industries, Ltd.,

Second digestion: Endoproteinase Asp-N (manufactured by catalog No. 129-02541)

rhird digestion: Trypsin-TPCK (manufactured by Worthington Boehringer-Mannheim Corp., catalog No. 1054 589) Biochemical, catalog No. 3740)

Peptide fragments obtained by each enzyme digestion were recovered, applied to a Wakosil-II 5C18 C18 eluted using a solvent A (0.05% TFA) and a solvent B inear gradient of from 1% B to 50% B at a flow rate of 0.25 peptide mappings were made (see Fig. 4). The resulting peptide ndustries, Ltd., 2.0 mm in diameter and 150 mm in length) and (isopropanol:acetonitrile = 7:3, 0.02% TFA), by 30 minutes of ragments were recovered and subjected to Edman degradation using a gas phase amino acid sequencer (PPSQ-2, manufactured by Shimadzu Corp.). Thereafter, each amino acid recovered in urn from the sequencer, whose N-terminal has been coupled chromatography by isocratic elution. The results are reverse phase column (manufactured by Wako Pure Chemical ml/min and at a column temperature of 30 c. In this way, with PTH, was identified using the C18 reverse phase column summarized in the following.

Amino acid sequences of peptide fragments obtained by the first digestion with lysyl endopeptidase

Fragment Amino acid sequence

(K)XYYESZ (X is A, S, G, M or Q and Z is E or K) (SEQ ID NO: 184) AP3

(K)XRAAZ (X is E or Q and Z is E or N) (SEQ ID NO: AP6

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(K)AGXCSG (cannot be totally identified) (SEQ ID NO: 186) AP7

NXPVPPACDPRIL (X is I, T OR S) (SEQ ID NO: 187) AP8

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(K)DSFLADVK (SEQ ID NO: 188)

(cannot be identified) (cannot be identified) **AP20** AP5

(cannot be identified) **AP21**

(cannot be identified) AP23

Amino acid sequences of peptide fragments obtained by the second digestion with endoproteinase Asp-N

Fragment Amino acid sequence

(cannot be identified)

ASP1

(cannot be identified) ASP2

(cannot be identified) **ASP6**

(cannot be identified) ASP11

Amino acid sequences of peptide fragments obtained by the third digestion with trypsin-TPCK

Fragment Amino acid secuence

(K or R)TLPTXAVP (SEQ ID NO: 189) TP2

(K or R)TLPTXAVP

In the above sequences, those shown in parentheses in N-termini are amino acid residues which can be deduced from the systematic enzyme digestion.

amino acid sequences < homology search> Analysis of the homology of obtained (2)

sequences are contained in a known reported protein or there is a protein having similar sequences, they were analyzed using sequence analyzing software Mac Vector (Kodak international Biotechnologies, Inc.). As for the information on known proteins or known genes, Entrez Release 6 data base In order to know if the thus obtained amino acid National Center for Biotechnology Information, National

Library of Medicine, National Institute of Health, USA, published on August 15, 1993) was used. Data bases included therein are as follows.

Entrez Release 6 database

VCBI-GenBank, August 15, 1993 (Release 78.0)

EMBL, July 15, 1993 (Release 35.0 plus updates)

DDBJ, July 15, 1993

SWISS-PROT, April, 1993 (Release 25.0)

PIR, June 30, 1993 (Release 37.0)

PDB, April, 1993

PRF, May, 1993

dbEST, July 15, 1993 (Release 1.10)

J.S. and European Patents

As the result, the sequence (K)DSFLADVK of AP12 a rat corticosteroid-binding globulin (CBG) precursor [PIR data completely coincided with an internal sequence KDSFLADVK of base accession No. A40066; Smith and Hammond; "Rat corticosteroid-binding globulin: primary structure and physiological conditions.", Mol. Endocrinol., (1989), 3, 420 messenger ribonucleic acid levels in the liver under different

Further detailed examination revealed that a sequence KQYYESE (SEQ ID NO: 193) having high similarity to AP3 is contained in the rat CBG amino acid sequence. In the rat CBG, sequences which correspond to those of AP12 and AP3 the sequence (K)XYYESZ (X is A, S, G, M or Q and Z is E or K) of are linked to each other thereby forming an internal amino acid sequence KDSFLADVKQYYESE (SEQ ID NO: 190).

fragments than AP12 and AP3, no protein or gene was found With regard to the amino acid sequences of other whose similarity could be taken into consideration.

<Example 3>

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Analysis of biological characteristics of TPO derived from blood plasma of thrombocytopenic rats

derived from blood presure of undimpositive rais
(1) In the rat CFU-MK assay system (liquid culture system)

As a typical example, Fig. 5 shows a dose response rom thrombocytopenic rat plasma (TPO-active fraction F2 naturation of megakaryocytes, namely increase in the cell (Topp et al., Blood, vol.76, pp.912 - 924, 1990), which is iar observable in vitro. Since such a morphological change was it is possible that this factor alone can stimulate proliferation and differentiation of CFU-MK, can produce curve in the case of the use of a TPO sample partially purified rom YMC Pack Protein-RP column described in the purification step (8) of Example 1-2). When the cultured cells were observed periodically under a microscope, generation and size, was recognized, probably together with increase in the change, process formation by megakaryocytes were found on the 4th day which was the final day of the culturing (they were hardly recognizable on the 3rd day). Such a process formation is called cytoplasmic process formation (Leven and Yee, Blood, vol.69, pp.1046 - 1052, 1987) or proplatelet process formation considered to be a platelet precursor structure further differentiated from mature megakaryocytes and is considered to be at the final stage of megakaryocyte differentiation so observed with a high frequency by using the TPO sample alone, number of megakaryocytes. As an especially significant mature megakaryocytes and can finally release platelets.

(2) In the colony assay system

When a TPO sample partially puritied from blood plasma of thrombocytopenic rats was examined by the colony assay system using unseparated rat bone marrow cells, cells of each of the separation/concentration steps or a Gplib/IIIa+CFU-MK fraction, the rat plasma-derived TPO stimulated formation of megakaryocyte colonies. In comparison with megakaryocyte colonies induced by other cytokine such as rat

IL-3, mouse GM-CSF or human EPO, the TPO-induced megakaryocyte colonies are characterized in that each colony consists of smaller numbers of megakaryocytes, but each megakaryocyte is larger in size, which means advanced maturity. In addition, since TPO produced no or few colonies of the other cell lineages, the Meg-CSF activity of TPO can be regarded as megakaryocyte-specific. On the basis of these facts, it is evident that TPO has different biological properties from those of other cytokines such as rat IL-3, mouse GM-CSF and human EPO and exerts unique Meg-CSF activity.

The TPO sample partially purified from blood plasma of thrombocytopenic rats also exerted its Meg-CSF activity on CD34+,DR+ cell fractions derived from human bone marrow cells or human cord blood cells and induced significant numbers of human megakaryocyte colonies, indicating that this factor has no species specificity.

<Example 4>

Specialization of rat TPO producing cells

(1) Screening of rat TPO producing organs

Screening and specialization of rat TPO-producing organs were carried out in order to ensure a mRNA source for use in the cloning of a rat TPO gene. Firstly, bone marrow, lungs, livers and spleens were taken periodically from rats rendered thrombocytopenic by P55 antibody administration, their cells (organ sections in the case of lungs and livers) were cultured and activities in the resulting culture supernatant were examined by the rat CFU-MK assay system. This initial attempt, however, did not yield distinctive results. In consequence, further attempt was made to culture hepatocytes prepared by collagenase perfusion from livers of thrombocytopenic rats induced by P55 antibody administration, taking into consideration a report which indicated a

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Biochemical and biological properties of the TPO-active proteins secreted from these 3 rat cell lines were examined in detall and compared with those of the rat plasmaderived TPO.

to recover the resulting culture supernatant. H4-II-E cells were suspended in Dulbecco's modified Eagle liquid culture iquid culture medium containing 10% FCS and transferred into After 3 days of culturing at 37 c in a 5% CO2 incubator, the 5% FCS and the culturing was continued for additional 3 days medium (glucose 4.5 g/l, to be referred to as "DMEM" McA-RH8994 cells were suspended in alpha-MEM(-) medium was exchanged with IMDM culture medium containing hereinafter) containing 10% FCS and transferred into a 175 cm² plastic tissue culture flask at $5 imes 10^5$ cells/flask. After 3 days of culturing at 37 c in a 5% CO2 incubator, the medium was exchanged with IMDM culture medium containing 5% FCS and the culturing was continued for additional 3 days to recover the resulting culture supernatant. Also, HTC cells FCS and transferred into a 175 cm² plastic tissue culture lask at 2.5 x 10⁵ cells/flask. After 3 days of culturing at IMDM culture medium containing 5% FCS and the culturing was a 175 cm² plastic tissue culture flask at 1 x 10⁶ cells/flask. were suspended in DMEM liquid culture medium containing 5% 37 c in a 5% CO2 incubator, the medium was exchanged with continued for additional 3 days to recover the resulting culture supernatant.

Using a 2 liter portion of the thus obtained culture supernatant from each of the 3 cell lines, partial purification of the cell line-derived TPO was carried out in accordance with the procedure for the purification of TPO from XRP described in Example 1-2. The following describes an outline of the results.

relationship between the liver and TPO production in rats (Siemensma et al., J. Lab. Clin. Med., vol.86, pp.817 - 833, 1975). When the resulting culture supernatant was applied to a WGA-Agarose column and then the adsorbed fraction was fractionated on a Vydac phenyl reverse phase column, exceedingly similar activity was found at the same position of the rat plasma-derived TPO activity in the rat CFU-MK assay system. Weak but the same activity was also found in culture supernatant of normal rat hepatocytes. These results strongly suggested that the liver would be one of the TPO-producing organs.

(2) Screening of rat TPO-producing cell lines

TPO-producing cell lines was carried out. Firstly, each of 20 rat liver-derived cell lines was cultured in respective same respective medium supplemented with 5% FCS, and the the procedure described in the above step (1) to examine existence of TPO production, the TPO activity was II-E cells (ATCC deposit No. CRL1548; Pitot et al., Nat. Cancer Inst. Monogr., vol.13, pp.229 - 245, 1964; purchased from Dainippon Pharmaceutical Co., Ltd.) and HTC cells (Thompson et al., Proc. Natl. Acad. Sci. USA, vol.56, pp.296 - 303, 1966; On the basis of the above results, screening of rat subculture liquid medium until the cells reached almost confluent stage, the culture medium was exchanged with the culturing, was continued for 3 days. When each of the resulting culture supernatant was partially purified in accordance with distinctively found in three rat hepatic parenchymal cellderived cell lines, namely McA-RH8994 cells (ATCC deposit No. ed. by Fishman and Sell, Academic Press, NY, pp.259 - 270, 1976; purchased from Dainippon Pharmaceutical Co., Ltd.), H4-CRL1602; Becker et al., "Oncodevelopmental Gene Expression" purchased from Dainippon Pharmaceutical Co., Ltd.).

(3) Detailed analysis of TPO activities in McA-RH8994 cells. H4-II-E cells and HTC cells



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using a Sephadex G-25 column. The resulting eluate was raction was concentrated and then fractionated using a about 6 times using an ultrafiltration apparatus and the medium was exchanged with 20 mM Tris-HCl buffer (pH 8.0) applied to a Q-Sepharose FF column, the column was washed with 20 mM Tris-HCI (pH 8.0) and then the adsorbed fraction was eluted with 20 mM Tris-HCI (pH 8.0) containing 175 mM NaCl. The thus eluted fraction was applied to a WGA-Agarose containing 0.2 M GlcNAc and 0.15 M NaCl. The resulting eluate everse phase Vydac Protein C4 column (manufactured by The Firstly, each culture supernatant was concentrated column, the column was washed with PBS and the adsorbed fraction was eluted with 20 mM sodium phosphate (pH 7.2) was applied to an TSK-gel AF-BLUE 650MH column, the column M NaCl and then the adsorbed fraction was eluted with 2 M sodium phosphate (pH 7.2) containing 1.5 M ammonium sulfate followed by 0.8 M ammonium sulfate containing 36 mM sodium phosphate, and then the adsorbed fraction was eluted with 20 and 15 cm in bed height). That is, the sample was applied to was washed with 20 mM sodium phosphate (pH 7.2) containing NaSCN. The resulting eluate was applied to a Phenyl-Sepharose 6 FF/LS column, the column was washed with 50 mM mM sodium phosphate (pH 7.2). The thus obtained adsorption Separations Group, catalog No. 214TP51015; 1 cm in diameter the C4 column which has been equilibrated in advance with 20% B and then elution was effected by 90 minutes of linear gradient of from 20% B to 40% B at a flow rate of 1 ml/min, using 0.1% TFA in the developing solvent A and 1-propanol containing 0.05% TFA in the developing solvent B. As the esult, each of the TPO-active proteins derived from these cell ines was eluted at a 1-propanol concentration of 30 to 43%.

When TPO activities in samples of each purification step were measured by the rat CFU-MK assay system, the results showed that TPO activities of the 3 cell

were measured by the rat CFU-MK assay system, TPO activities of the 3 cell lines and the XRP-derived TPO were found in the same peak area. When colony assay of TPO-active fractions elution patterns of Meg-CSF activity closely resembled those of TPO activity measured by the rat CFU-MK assay system (see Table 3). In addition, similar to the case of the XRP-derived from the reverse phase column was carried out using TPO, each of TPO activities of the 3 cell lines formed no or MK assay system are shown in Table 2. In addition, when TPO activities in the eluates from the final reverse phase column nonadherent cells obtained in the adherence depletion step for the separation and concentration of rat GpIIb/IIIa+ CFU-MK, ines behaved with the similar patterns to those of the XRP-Relative activity, activity yield and the like at each purification step measured by the rat CFUew colonies of the other cell lineages. derived TPO (cf. Example 1-2).

Each of the pooled active fractions from the reverse phase column was subjected to SDS-PAGE in accordance with the procedure described in Example 1-2. When derived TPO and H4-II-E cell-derived TPO were found to be respectively, and the HTC cell-derived TPO showed apparent protein was extracted from the resulting gel and TPO activity was measured by the rat CFU-MK assay system, apparent 17,000 to 22,000, 33,000 to 39,000 and 31,000 to 38,000, molecular weights of the XRP-derived TPO, McA-RH8994 cellmolecular weights of 17,000 to 22,000 and 28,000 to 35,000.

Thus, these results revealed that the TPO proteins produced by McA-RH8994 cells, H4-II-E cells and HTC cells are properties, though their biochemical properties are slightly different from one another in terms of apparent molecular blood may be a product of partial digestion at its specific or equivalent to the XRP-derived TPO in terms of their biological weight. A remarkable finding in the present invention is possibility that a TPO-active protein molecule contained

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from the production cells. It also suggests possible existence of TPO genes (mRNA and cDNA) having various lengths. irregular site in its production cells or after its secretion

Table 2 Purification of TPO from rat cell lines

,	•								
6 FF/LS Vydac Protein C4	62.0	_		SL.0	_	_	11.0		.
Phenyl-Sepharose	L'\$1	200	0\$£L	1.61	2500	35750	4.11	0 <i>L</i> I	0761
TSK-gel AF-Blue	0.8	15200	92200	2.6	00 <i>SL</i>	00069	4.2	051	018
92018gA-ADW	0.97	1320	105600	132.0	220	33000	0.08	06	7200
Q-Sepharose FF	£261	50	39460	3115	١٤٠	08997	1831	51	27320
Sephadex G25	4854	91	.081 <i>LL</i>	8549	15	00\$ <i>LL</i>	1101	ς	20020
Culture supernatant	(an) 4806	8	38420	09 <i>L</i> Տ	ς	28800	(8tu)	S	20440
Z(cl)	$_{1*}dL$	RA*²	C+AT	ЧL	AЯ	AT	ФŢ	VИ	ΛT
Cell Line	McA-	₽ <u>6</u> 68H8	•		DTH		·II-ÞH	E	

, total protein; 2* , relative activity; 3* , total activity

Þ	0	Þ	O	%0.Eb - 9.Ob			
28	ς	12	7	38.8 - 40.9%			
103	183	717	0L	%8.8E - 7.8E			
230	167	790	<i>L</i> 9	34.7 - 36.7%			
05	61	135	0	32.6 - 34.7%			
81	Þ	33	0	30.0 - 32.6%			
səinoloə	colonies	colonies	soinoloo	propanol			
H4-II-B	DTH	McA-RH8994	ЯЯХ	noimla			
Table 3 Rat megakaryocyte colony formation by rat TPO (reverse phaseC4 column fraction)							

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Construction of expression vector (pEF18S) for cDNA library

A vector into which a cDNA fragment can be ntegrated easily and which can exert a high expression efficiency of the cloned cDNA was constructed for use in the cloning and expression of TPO cDNA. That is, an expression vector pEF18S was constructed from an expression vector digested in advance with Hindill and EcoRI. Cells of a commercial host strain, Competent High E. coli DH5 (manufactured by Toyobo Co., Ltd.; a competent E. coli strain vol.166, pp.557 - 580, 1983), were transformed with the thus constructed vector. After culturing the transformed cells, 12 colonies were selected at random and plasmid DNA was purified from each colony. A total of 10 clones containing a plasmid of interest (pEF18S) were obtained by analyzing pME18S by replacing the SRlpha promoter with an elongation factor 1α (EF1α) promoter which is known as a high expression promoter (cf. Fig. 6). The promoter of elongation factor 1α was obtained by partially digesting 1 μg of an expression vector pEF-BOS (Mizushima et al., Nucleic Acids Res., vol.18, subjecting the digest to 2% agarose gel (manufactured by FMC BioProducts) electrophoresis to isolate a DNA fragment of about 1,200 bp and purifying the DNA fragment using a Prep-A-Bio-Rad Laboratories, Inc.; a kit for use in the selective purification of DNA which is effected by porous silica matrix-aided DNA adsorption, developed on the basis of the procedure reported by Willis et al. in Bio Techniques, vol.9, pp.92-99, 1990). A 100 ng portion of the thus obtained DNA fragment was ligated with 50 ng of an expression vector pME18S (Liu et al., Proc. Natl. Acad. Sci. USA, vol.90, pp.8957 - 8961, 1993) which has been prepared by a modified method of Hanahan et al., J. Mol. Biol., p.5322, 1990) with restriction enzymes Hindlll and EcoRI Gene DNA purification kit (manufactured by

Thereafter, a clone selected therefrom was cultured to prepare estriction enzyme digestion patterns of these DNA molecules. a large quantity of the plasmid DNA.

manufactured by FMC BioProducts) electrophoresis, and the aboratory Press, 1989). That is, the clone pEF18S obtained glucose, 0.5% lysozyme). To this were added 8 ml of 0.2 N he same volume of isopropanol and then centrifuged. The pH 7.5, 1 mM EDTA) and treated with RNase and then with the resulting pellet was again dissolved in TE solution to which were subsequently added NaCl and polyethylene glycol respectively. After centrifugation, the pellet was dissolved in IE solution and precipitated with ethanol. In this way, about 300 µg of the plasmid DNA was obtained. A 100 µg portion of the DNA was completely digested with restriction enzymes EcoRi and Notl and subjected to 0.8% agarose gel thus recovered vector fragments were purified using a Prep-A-Gene DNA purification kit (manufactured by Blo-Rad basically in accordance with the procedure described in Nolecular Cloning (Sambrook et al., Cold Spring Harbor above was cultured overnight at 37 c in 50 ml of an LB medium containing 50 µg/ml of ampicillin, and the resulting cells ysozyme solution (25 mM Tris-HCI, pH 8, 10 mM EDTA, 50 mM VaOH/1% SDS solution and then 6 ml of 3 M potassium/5 M acetate solution to suspend the cells thoroughly. After centrifugation of the suspension, the resulting supernatant fluid was treated with phenol/chloroform (1:1), mixed with shenol/chloroform (1:1), followed by ethanol precipitation. 3,000 to their final concentrations of 0.63 M and 7.5%, Laboratories, Inc.) to obtain about 55 µg of plasmid DNA which Purification of the plasmid DNA was carried out (1% Bacto-tryptone, 0.5% Bacto-yeast extract, 0.5% NaCl) collected by centrifugation were suspended in 4 ml of TEGresulting pellet was dissolved in TE solution (10 mM Tris-HCI, vas used in the subsequent cDNA library construction.

<Example 6>

Purification of mRNA from McA-RH8994 cells

McA-RH8994 cells which showed relatively high activity in the colony assay of Example 4 were selected as the material for use in the rat TPO cDNA cloning and subjected to he following experiments.

about 20 ml was gently layered over the cushion in such a nours of centrifugation at 25,000 rpm and at 20°c. The esulting pellet was washed twice with a small amount of 80% bhenol/chloroform (1:1) and then subjected to ethanol precipitation with 1/10 volume of 3 M sodium acetate and 2.5 disruption, was subjected to about 20 repetitions of suction/discharge with a 20 ml syringe equipped with a 18G needle and then 21G needle until the mixture became almost Ise in Beckman SW28 rotor, and the aforementioned mixture of nanner that the layers were not disturbed and the tube was almost filled. The thus prepared tube was subjected to 20 ethanol, dissolved in TE solution, extracted with Isolation of total RNA was carried out basically in accordance with the procedure described in Molecular Cloning That is, McA-RH8994 cells were grown to confluence in 15 culture dishes of 90 mm in diameter. After removing the iquid medium, cells in each dish were thoroughly suspended in 3.8 ml of a 5 M guanidine solution (5 M guanidine thiocyanate, 5 mM sodium citrate (pH 7.0), 0.1 M β-mercaptoethanol, 0.5% sodium sarcosylsulfate), and the resulting suspensions in all dishes were collected in a single tube and the total volume esulting mixture, which became viscous due to cell non-viscous. A 18 ml portion of 5.7 M CsCl-0.1 M EDTA (pH 7.5) vas used as a cushion in a polyallomer centrifugation tube for Sambrook et al., Cold Spring Harbor Laboratory Press, 1989). vas adjusted to 20 ml with the guanidine solution.

volumes of ethanol. In this way, about 2.5 mg of total RNA was

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by Japan Synthetic Rubber/Nippon Roche; oligo dT molecules vas carried out using Oligotex TM-dT30 (Super) (manufactured are immobilized on latex particles by covalent bonding, and purification of poly (A)+ RNA can be attained similar to use of As the result, 20 µg of poly (A)+ RNA was obtained from about Purification of poly (A)+ RNA from the total RNA an oligo dT column which is generally used for such purpose). obtained from about 108 cells. 500 µg of the total RNA.

<Example 7>

Construction of rat cDNA library

site at its 5'-end and a Notl recognition site at the 3'-end was synthesized from 5 µg of the poly (A)+ RNA obtained in (manufactured by Pharmacia; a set of a primer 5'-AACTGGAAGAATTCGCGGCCGCAGGAA(T)₁₈-3' (SEQ ID NO:15) Double-stranded cDNA having an EcoRI recognition Example 6, using a Time SaverTM cDNA synthesis kit manufactured by Pharmacia; a cDNA synthesis kit based on the containing a Notl recognition sequence for use in the synthesis and 5'-CTCGTGCCG-3' (SEQ ID NO:17) for use in the addition of Not! and transformed into 8.4 ml of the aforementioned Competent High E. coli DH5 (manufactured by Toyobo Co., Ltd.) modified method of Okayama and Berg, Mol. Cell. Biol., vol.2, pp.161 - 170, 1982) and DIRECTIONAL CLONING TOOLBOX of cDNA and adapters 5'-AATTCGGCACGAG-3' (SEQ ID NO:16) an EcoRI recognition sequence). The thus synthesized cDNA Example 5) which had been digested in advance with EcoRI and was ligated with 1.2 µg of the expression vector pEF18S (cf. is the result, 5.3 x 10⁵ transformants were obtained.

<Example 8>

Preparation (cloning) of rat TPO cDNA fragment by PCR

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RH8994 cDNA library plasmids were purified from the ip100 (manufactured by DIAGEN; an anion exchange silica The 530,000 clones of the McA-RH8994 cDNA ibrary constructed in Example 7 were cultured overnight in 50 ml of LB medium containing 50 µg/ml of ampicillin, and McAresulting cells collected by centrifugation, using QIAGENcolumn for DNA purification use). About 200 µg of plasmid ONA was obtained.

nucleotide primers EF1a-1 and EF1a-2, which correspond to he first intron of the human elongation factor 1α of the he p-cyanoethylamidite method) and their purification was synthetic DNA having trityl groups). Each of the thus synthesized and purified DNA molecules was dissolved in TE solution to a final concentration of 50 µM and stored at -20 °C Two anti-sense nucleotide primers AP8-1R and AP8-2R, which correspond to the amino acid sequence of the peptide fragment AP8 described in Example 2, and two sense plasmid vector pEF-18S (cf. Fig. 6) used for the preparation of Synthesis of these primers was effected making use of a 394DNA/RNA Synthesizer manufactured by Applied Biosystems; a synthesizer based on effected by the use of an OPC column for synthetic DNA purification use (manufactured by Applied Biosystems; a reverse phase silica gel column for use in the purification of until its use. Synthetic oligonucleotides used in the following procedure were synthesized and purified in the same manner. cDNA library, were synthesized.

The primers AP8-1R and AP8-2R are mixed primers consisting of 17 continued nucleotides in which deoxyinosine is incorporated, as reported by Takahashi et al. (Proc. Natl. Acad. Sci. USA, vol.82, pp.1931 - 1935, 1985).

the nucleotide sequences of positions 1491 to 1512 and 1513 and 20 nucleotides, respectively, were synthesized based on The primers EF1a-1 and EF1a-2 consisting of 21

to 1532 of the genomic sequence reported by Uetsuki et al. (J. *Biol. Chem.*, vol.264, pp.5791 - 5798, 1989)

(SEQ ID NO:18) (SEQ ID NO:19) ID NO:20) (SEQ (SEQ ID NO:21) (SEQ ID NO:22) (SEQ ID NO:23) SEQ ID NO:24) (SEQ ID NO:25) AP8: Ile Pro Val Pro Pro Ala Cys Asp Pro Arg Leu Leu GAI GA-5' AP8-2R: 3'-GGG GGG CGG ACG CTG GG-5' 4 AP8-1R: 3'-ACG CTG GGG GCI

SF1a-2: 5'-CCT CAG ACA GTG GTT CAA AG-3' (SEQ ID NO:30) FE1a-1: 5'-GGA TCT TGG TTC ATT CTC AAG-3' (SEQ ID NO:29)

(SEQ ID NO:26) SEQ ID NO:27) SEQ ID NO:28)

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ncubation at 72.c for 7 minutes). In order to improve Using 3 µg of the McA-RH8994 cDNA library olasmid as a template, AP8-1R (500 pmol) and EF1a-1 (100 AmpliTaq TM DNA Polymerase (manufactured by Takara Shuzo Co., Ltd.; a set of a thermostable Taql polymerase, a reaction buffer and dNTP for use in PCR), PCR was carried out by specificity of the thus amplified DNA fragments, further PCR pmol) as primers and GeneAmpTM PCR Reagent Kit with GeneAmpTM PCR system 9600 (manufactured by Perkin-Elmer; a thermal cycker for PCR) (100 µl volume; heating at 95 c for denaturation at 95 c for 1 minute, annealing at 40 c for 1 minute and synthesis at 72°c for 1 minute, and final 2 minutes, a total of 35 cycles, each cycle consisting of vas carried out (100 µl volume; heating at 95·c for 2 minutes,

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synthesis at 72 c for 1 minute, and final incubation at 72 c a total of 35 cycles, each cycle consisting of denaturation at 95.c for 1 minute, annealing at 45.c for 1 minute and for 7 minutes), using 1 µl of the thus obtained PCR reaction solution as a template, and EF1a-2 (100 pmol) and AP8-2R (500 pmol) as primers.

The thus obtained reaction solution was subjected the main product of the PCR which was subsequently purified Since the thermostable Cycle Sequening Kit (manufactured by Applied Biosystems; a to 2% agarose gel (manufactured by FMC BioProducts) electrophoresis to isolate a DNA fragment of about 330 bp as naking use of the aforementioned Prep-A-Gene DNA purification kit (manufactured by Bio-Rad Laboratories, Inc.). Using T4 DNA ligase (manufactured by Life Technologies), the hus purified DNA fragment was subcloned into pCRTMII vector (a vector for the use of TA cloning of PCR products, polymerase to be used in PCR has a terminal transferase activity, the amplified DNA fragment is directly subcloned into the pCRTMI vector which has a 5'-dT cohesive end, making use of the enzyme's property to add one deoxyadenylic acid to the 3'-end of the PCR-amplified DNA. Plasmid DNA molecules were purified from 28 clones selected at random from the resulting clones using QIAGEN-tip100 (manufactured by DIAGEN) and their nucleotide sequences were determined by a 373A DNA sequencer (a fluorescence sequencer, manufactured by Applied Biosystems), making use of a Taq Dye DeoxyTM Terminater tit for use in PCR-aided nucleotide sequence determination using fluorescent dyes, based on the dideoxy method of Sanger et al., Proc. Natl. Acad. Sci. USA, vol.74, pp.5463 - 5467, 1977). manufactured by Invitrogen).

residues, (Ile/Thr/Ser)-Val-Pro, of the amino acid was selected from the thus obtained DNA fragments and its When a DNA fragment which encodes three amino sequence of AP8 shown above, adjacent to the AP8-2R primer,

Kozak (Kozak, M., Cell, vol.44, pp.238 - 292, 1986), it seems hat this cDNA fragment contains a translation initiation region, coding for the N-terminal of rat TPO protein. This Position 173 to 175 of this cDNA fragment encoded nethionine, and the coding frame coincided with the amino acid frame of AP8. Since the sequence interposing the position 173-175 of the DNA fragment coincided with the sequence of ragment excluding the vector sequence, and an amino acid cDNA fragment was named A1. Nucleotide sequence of the A1 sequence deduced therefrom are shown in the Sequence Listing whole length was sequenced, it contained cDNA of 261 bp. SEQ ID NO: 1) attached hereto.

<Example 9>

Screening of rat TPO cDNA clone by PCR

pools of about 10,000 clones, cultured overnight in 1 ml of aforementioned LB medium containing 50 µg/ml of ampicillin nanufactured by Kurabo Industries, Ltd.; an automatic plasmid DNA extraction apparatus based on a modified method of the alkali SDS method disclosed in Molecular Cloning, Sambrook et from this, the following two oligonucleotides were and then subjected to plasmid DNA extraction using an The aforementioned cDNA library was divided into automatic plasmid isolation apparatus PI-100 (VER-3.0, al., Cold Spring Harbor Laboratory Press, 1989). Separately synthesized based on the cDNA fragment A1 and purified.

5' CGAGGGTGTACCTGGGTCCTG 3' (SEQ ID NO:31) (a sense sequence of positions 1 to 17 of the sequence of SEQ ID NO: 1; CGAG represents an adapter sequence)

5' CAGAGTTAGTCTTGCGGTGAG 3' (SEQ ID NO:32) (an anti-sense sequence of positions 212 to 232 of the sequence of SEQ ID NO: Using 1/30 volume of each plasmid DNA sample above as a template, the thus synthesized obtained

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with AmpliTaqTM DNA Polymerase (manufactured by Takara Shuzo Co., Ltd.), PCR was carried out by GeneAmpTM PCR System 9600 (manufactured by PERKIN-ELMER) (a total of 30 oligonucleotides as primers and GeneAmpTM PCR Reagent Kit cycles, each cycle consisting of denaturation at 94.c for 30 seconds, annealing at 66 c for 30 seconds and synthesis at 72.c for 1 minute). As the result, a specific band of 236 bp was detected in 3 of the 100 pools examined. When one of these 3 pools was divided into sub-pools of about 900 clones to extract plasmid DNA and PCR was carried out in the same manner, a specific band was detected in 3 of the 100 sub-pools ested. When one of these sub-pools was further divided into bools of 40 clones and the same screening process was carried One of the candidate pools was cultured on an LB plate (LB medium containing 15% agar) supplemented with 50 µg/ml of ampicitlin and each of the thus formed colonies was subjected to plasmid DNA extraction and PCR in the same manner. As the esult, a positive band was detected in 2 of the 100 clones out, the specific band was found in 3 of the 100 pools tested. examined.

<Example 10>

Sequencing of rat TPO cDNA

Molecular Cloning (Sambrook et al., Cold Spring Harbor Laboratory Press, 1989). Each of the two clones obtained in basically in accordance with the procedure described in Example 9 were cultured overnight in 50 ml of LB medium containing 50 µg/ml of ampicillin, and about 300 µg of plasmid DNA was obtained after purification in the same Purification of plasmid DNA was carried out nanner as described in Example 6.

The thus obtained plasmid DNA was subjected to sequencing in the same manner as described in Example 8 using the Taq Dye DeoxyTM Terminater Cycle Sequening Kit

manufactured by Applied Biosystems) to determine complete coincided with each other, showing that they are the same clone. One of these clones was selected, and the plasmid carried by this clone was named pEF18S-A2α. Its nucleotide sequence and an amino acid sequence deduced therefrom are As the esult, nucleotide sequences of the two clones completely nucleotide sequence containing the A1 fragment. shown in the Sequence Listing (SEQ ID NO: 2).

of the 172 bp 5' non-translation region encodes an amino acid sequence rich in hydrophobic amino acids which is presumed to be a protein secretion signal sequence consisting of 21 amino acids starting with methionine. This protein contains 126 The 1624-1629 nucleotide sequence located closely to the terminal of the 3' non-translation sequence is different from a The sequence shown in the Sequence Listing (SEQ ID VO: 2) has distinct characteristics. The sequence downstream amino acid residues, and a 3' non-translation sequence of 1,025 nucleotides and a poly A tail following the termination codon (TAA). This protein contains a sequence which corresponds to the amino acid sequence AP8 analyzed in Example 2 (amino acid numbers 1 to 12 in SEQ ID NO: 2), but does not contain a consensus sequence for N-glycosylation. consensus sequence but seems to be a potential polyadenylation sequence.

The vector pEF18S-A2a carried by an E. coli strain DH5 has been deposited by the present inventors on February 14, 1994, under the deposit No. FERM BP-4565, in National nstitute of Bioscience and Human Technology, Agency of ndustrial Science and Technology, Ministry of International frade and Industry, Japan.

Example 11>

sells and confirmation of TPO activity Expression of rat TPO cDNA in COS

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extensively dialyzed against IMDM culture medium and evaluated by the rat CFU-MK assay system. TPO activity was ound in a dose-dependent fashion in the culture supernatant of COS 1 cells in which the plasmid pEF18S-A2 α was expressed negakaryocytes formed elongated cytoplasmic processes on 4 The thus obtained culture supernatant was (Fig. 7). Similar to the case of XRP-derived TPO, many

supernatant.

days of culture. By contrast, TPO activity was not found in the culture supernatant of COS 1 cells to which the plasmid In the M-07e assay system, M-07e cell proliferation enhancing activity was also found in a dose dependent fashion in the culture supernatant of but not in the culture supernatant of COS 1 cells in which the plasmid pEF18S alone was expressed. These results demonstrated that A2a contains a cDNA which encodes a COS 1 cells in which the plasmid pEF18S-A2 α was expressed, DEF18S alone was transfected (Fig. 7). protein having a TPO activity.

in order to examine for the ability of this TPO activity to transfected with the plasmid pEF18S-A2 α were cultured for 3 promote platelet production in vivo. Firstly, COS 1 cells days in a serum free culture medium which has been supplemented with 0.2 mg of BSA, thereby obtaining about 5.8 APMSF was added to the serum-free culture supernatant to the using a 0.22 µm filter. To 5,793 ml of the resulting filtrate relative activity, 1,000; total activity, 1,326,000) was added obtaining 5,849 ml of solution containing 0.822 M NaCl. The Next, a partially purified TPO sample was prepared iters of serum free culture supernatant. Protease inhibitor pfinal concentration of 1 mM, and the mixture was filtered 0.85 moles of NaCl for 1000 ml (288 g in total), thereby to a TSK-gel AF-BLUE 650 MH column (manufactured by Tosoh Corp., catalog No. 08705; 5 cm in diameter and 6 cm in bed completion of the sample application, about 7,900 ml of eluate bassed through the column when eluted with 20 mM sodium and concentrated using an ultrafiltration unit (Omega (protein concentration, 0.229 mg/ml; total protein, 1,326 mg; neight) which has been equilibrated in advance with 20 mM shosphate (pH 7.2) containing 1 M NaCl. This eluate was pooled thus prepared solution was applied, at a flow rate of 7 ml/min, Ultrasette, nominal molecular weight cutoff of 8,000; sodium phosphate (pH 7.2) containing 1 M NaCl.

raction F1 (460 ml; protein concentration, 2.11 mg/ml; total protein, 973 mg; relative activity, 16.3). Next, the elution

manufactured by Filtron), thereby obtaining a passed-through

solution was changed to 2 M NaSCN, and the thus eluted TSK-Jel AF-BLUE 650MH-adsorbed TPO-active fraction F2 (2840 nl) was concentrated to 6.81 ml using an ultrafiltration unit with YM 3 membrane (76 mm in diameter, manufactured by

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concentration, 4.74 mg/ml; total protein, 7.82 mg; relative activity, 71,400; total activity, 558,600)

subcutaneously administered for consecutive 5 days (474 µg (100 µl)/mouse/day) to ICR male mice (9 weeks old) whose platelet numbers had been measured on the day before the As a control, BSA (200 μg (100 ourified TPO (100 µl/mouse/day) was administered in the ul)/mouse/day) or a buffer used as the solvent of the partially blood was collected from the heart of each mouse to measure mice, an increase in platelet counts by a factor of about 2.14 was observed in comparison with platelet counts before the same manner. On the next day of the final administration, platelet counts using a hemocytometer (F800, manufactured by In the partially purified TPO-administered administration. When compared with the control groups, increased platelet counts in the partially purified TPOadministered mice were about 1.74 times higher than those in the BSA-administered mice and about 1.90 higher than those in he buffer-administered mice. These results demonstrated since the amount of immunosuppressive acidic protein (IAP), known as a mouse acute phase protein, was not elevated in the that TPO promotes platelet production in vivo. In addition, partially purified TPO-administered mice, it was strongly suggested that TPO is different from IL-6 and IL-11 with purified TPO sample egard to the induction of acute phase protein. partially oa Iyo Denshi). administration.

(manufactured by Pharmacia Biotech, catalog No. 17-1071-01;

2.6 cm in diameter and 60 cm in bed height) and developed with 20 mM sodium acetate (pH 5.5) containing 50 mM NaCl at a low rate of 1 ml/min. After commencement of the development, TPO activity was found in eluates in the range of

from 194 to 260 ml after application. These eluates were pooled to obtain a Superdex 200 pg TPO-active fraction F2 (66 ml; protein concentration, 0.112 mg/ml; total protein, 7.41 mg;

Amicon Corp.). Total protein in this TPO-active fraction F2 was 12.5 mg, and protein yield of F2 at this step was 0.62%. Relative activity of TPO was calculated to be 240. Next, the F2 was applied to a HiLoad 26/60 Superdex 200 pg

<Example 12>

rate of 1 ml/min to a strong cation exchange column RESOURCE

S (manufactured by Pharmacia Biotech, catalog No. 17-1178-01; 0.64 cm in diameter and 3 cm in bed height) which has been

equilibrated in advance with 100% A. Thereafter, elution was carried out at a flow rate of 0.3 ml/min with 40 minutes of a

inear gradient of from 100% A to 100% B. When assayed, the TPO activity was detected in a broad range of eluates (from 5% B to 32% B). These TPO-active eluates were pooled and concentrated using an ultrafiltration unit with YM 3 membrane

(25 mm in diameter, manufactured by Amicon Corp.) to obtain a

RESOURCE S TPO-active fraction F2 (1.65 ml; protein

buffer A (20 mM sodium acetate, pH 5.5) and buffer B (20 mM sodium phosphate, pH 7.2, containing 500 mM NaCI) were prepared, and the TPO-active fraction was applied at a flow

relative activity, 142,860; total activity, 1,058,600). Next,

Detection of TPO mRNA in rat tissues

in the rat body, RNA was extracted from various rat tissues. A total of 6 rats were subjected to X-ray irradiation in the same In order to locate rat TPO mRNA expressing tissue manner as described in Example 1, and various tissues (brain, thymus, lung, liver, heart, spleen, small intestine, kidney,

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the 11th to 14th day after the X-ray irradiation and immediately frozen in liquid nitrogen. Extraction of total RNA was effected by the use of an RNA isolation reagent ISOGEN Amount of ISOGEN to be used was defermined according the instruments MFG. Co., LTD) until complete disintegration of the issue was attained (approximately 45 to 60 seconds at 10,000 rmp). The tissue homogenate was then subjected to total RNA extraction making use of a procedure based on the 1.1 to 5.6 mg of total RNA was obtained from respective testis and bone marrow cells) were excised from the rats on weight of each frozen tissue sample, and the reagent-added lissue sample was treated with a homogenizer (Physcotron R NS-60, manufactured by NITI-ON Medical & Physical acid guanidium phenol chloroform method of Chomczynski et al. (Anal. Biochem., vol.162, pp.156 - 159, 1987). As the result, (manufactured by Wako Pure Cher I Industries, Ltd.) issues.

Making use of Oligotex^{T M}-dT30 (Super) manufactured by Japan Synthetic Rubber/Nippon Roche), 20 µg of poly (A)+ RNA was purified from about 500 µg of total RNA.

Using random primer, first strand of cDNA was tissue. That is, 1 µg of the poly (A)+ RNA was dissolved in 10 ul of sterile water, incubated at 70°c for 15 minutes and then (Boehringer-Mannheim Corp.), 50 mM of Tris-HCl (pH 8.3), 75 mM of KCI, 3 mM of MgCl₂ and 200 U of Super ScriptTM II (a thus prepared solution (total volume, 20 µl) was incubated at synthesized from 1 µg of the poly (A)+ RNA obtained from each rapidly cooled down. To this were added 75 pmoles of random everse transcriptase manufactured by Life Technologies). The 37 c for 1 hour, and the resulting reaction solution was ncubated at 70°c for 10 minutes to inactivate the enzyme and primer (Takara Shuzo Co., Ltd.), 10 U of RNase inhibitor stored at -20°c until its use.

rat TPO cDNA sequence obtained in Example 10. Sequences of New primers for PCR were synthesized based on

TPO-I: 5-CCT GTC CTG CTG CCT GCT GTG-3' (SEQ ID NO:33) he thus synthesized primers are as follows. positions 347 to 367 in SEQ ID NO: 2)

anti-sense primer corresponding to positions 1005 to 1025 in rpo-n: 5-tga agt tcg tct cca aca atc-3' (SEQ ID NO:34) SEQ ID NO: 2)

results indicate that expression of TPO mRNA in rat occurs in be judged. When possible existence of similar expression mode in the human body and the results of Example 4 are taken Using 1/10 volume of each of the synthesized cDNA solution as a template, 1 µM each of the thus synthesized TPO-I and rTPO-N as primers and GeneAmpTM PCR Reagent Kit with AmpliTaqTM DNA Polymerase (manufactured by Takara PERKIN-ELMER) and heating at 95 c for 2 minutes, repeating a 95.c for 1 minute, annealing at 57.c for 1 minute and incubation at 72.c for 7 minutes. When each of the thus obtained reaction solutions was subjected to electrophoresis the amplified bands were examined, bands that seemed to be specific for TPO mRNA were found on gels resulting from the brain, the liver, the small intestines and the kidney. These tissues of these organs, though its expression quantity cannot together into consideration, it seems that the liver is an appropriate starting material for the acquisition of human TPO Shuzo Co., Ltd.), PCR was carried out in a volume of 100 µl, making use of GeneAmpTM PCR System 9600 (manufactured by total of 30 cycles, each cycle consisting of denaturation at synthesis at 72.c for 1 minute, followed by the final using 2% agarose gel (manufactured by FMC BioProducts) and

<Example 13>

Construction of human normal liver-derived cDNA library

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synthesized from 5 µg of commercial normal human liverextracted based on the acid guanidium phenol chloroform CLONING TOOLBOX (manufactured by Pharmacia). The thus synthesized cDNA was ligated with 1.2 μg of the selected as the starting material for use in the cloning of human TPO cDNA. In the same manner as described in Example 7, double-stranded cDNA having an EcoRI recognition site on its 5'-end and a Notl recognition site on its 3'-end was derived poly (A)+ RNA (manufactured by Clontech; a product method of Chomczynski et al.), using a Time SaverTM cDNA aforementioned expression vector pEF18S which had been Based on the results of Example 12, the liver was synthesis kit (manufactured by Pharmacia) and DIRECTIONAL digested in advance with EcoRI and NotI and transformed into 8.4 ml of the aforementioned Competent High E. coli DH5 (manufactured by Toyobo Co., Ltd.) As the result, 1.2 imes 10⁶ transformants were obtained.

<Example 14>

Preparation (cloning) of human TPO cDNA fragment by PCR

Using random primers, first strand of cDNA was ug of the poly (A)+ RNA was dissolved in 10 µl of sterile Mannheim Corp.), 50 mM of Tris-HCl (pH 8.3), 75 mM of KCl, 3 Script TM II (manufactured by Life Technologies). The thus prepared solution (total volume, 20 µl) was incubated at 37·c derived poly (A)+ RNA (manufactured by Clontech). That is, 1 cooled down. To this were added 75 pmoles of random primers mM of MgCl₂ and 200 U of a reverse transcriptase Super synthesized from 1 µg of the commercial normal human liver-(Takara Shuzo Co., Ltd.), 10 U of RNase inhibitor (Boehringerfor 1 hour, and the resulting reaction solution was incubated water, incubated at 70°c for 15 minutes and then rapidly

at 70 c for 10 minutes to inactivate the enzyme and stored at -20 c until its use.

Primers for PCR use were synthesized based on the rat TPO cDNA sequence (SEQ ID NO: 2). Sequences of the thus synthesized primers are as follows.

TPO-AIN: 5-ATG GAG CTG ACT GAT TTG CTC-3' (SEQ ID NO:35)

(positions 173 to 193 in SEQ ID NO: 2)

(anti-sense primer corresponding to positions 1005 to 1025 in rtpo-n: 5'-TGA AGT TCG TCT CCA ACA ATC-3' (SEQ ID NO:36) SEQ ID NO: 2)

solution as a template, 1 µM each of the thus synthesized rTPO-AIN and rTPO-N as primers and GeneAmpTM PCR Reagent Using 1/10 volume of the synthesized cDNA Kit with AmpliTaqTM DNA Polymerase (manufactured by Takara Shuzo Co., Ltd.), PCR was carried out with a volume of 100 μl, making use of GeneAmpTM PCR System 9600 (manufactured by PERKIN-ELMER) and heating at 95 c for 2 minutes, repeating a total of 35 cycles, each cycle consisting of denaturation at 95.c for 1 minute, annealing at 40.c for 1 minute and synthesis at 72 c for 1 minute, followed by the final incubation at 72.c for 7 minutes.

directly determined by a 373A DNA sequencer (manufactured excluding the primer moiety and an amino acid sequence The thus obtained reaction solution was subjected electrophoresis to isolate a DNA fragment of about 620 bp as the main product of the PCR which was subsequently purified making use of the aforementioned Prep-A-Gene DNA Nucleotide sequence of the thus purified DNA fragment was by Applied Biosystems) making use of the aforementioned Tag Dye DeoxyTM Terminater Cycle Sequening Kit (manufactured by Applied Biosystems). The thus determined nucleotide sequence to 2% agarose gel (manufactured by FMC BioProducts) purification kit (manufactured by Bio-Rad Laboratories, Inc.).

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deduced therefrom are shown in the Sequence Listing (SEQ ID NO: 3).

CDNA showed a homology of 86% with the rat cDNA nucleotide sequence, indicating that this DNA fragment encodes a portion sequences, has a length of 580 bp. When compared, the human This DNA fragment, excluding the of human TPO cDNA.

<Example 15>

Screening of human TPO cDNA clone by PCB

Human TPO-corresponding primers for PCR were Sequences of the thus synthesized based on SEQ ID NO: 3. synthesized primers are as follows.

htpo-1: 5-ttg tga cct ccg agt cct cag-3' (SEQ ID NO:37)

(anti-sense primer corresponding to positions 479 to 499 in hTPO-J: 5'-TGA CGC AGA GGG TGG ACC CTC-3' (SEQ ID NO:38) positions 60 to 80 in SEQ ID NO: 3) SEQ ID NO: 3)

. The human cDNA library constructed in Example 13 about 100,000 clones), cultured overnight in 1 ml of the and then subjected to plasmid DNA extraction using an manufactured by Kurabo Industries, Ltd.). The thus extracted was amplified and divided into pools (each pool containing aforementioned LB medium containing 50 µg/ml of ampicillin automatic plasmid isolation apparatus PI-100 (VER-3.0, DNA was dissolved in TE solution.

Using 5% of the extracted DNA sample as a ELMER) (a total of 35 cycles, each cycle consisting of (hTPO-I and hTPO-J) as primers and GeneAmpTM PCR Reagent Kit with AmpliTaqTM DNA Polymerase (manufactured by Takara denaturation at 95 c for 1 minute, annealing at 59 c for 1 template, 1 µM each of the synthesized oligonucleotides Shuzo Co., Ltd.), PCR was carried out in 20 µl volume by GeneAmpTM PCR System 9600 (manufactured by PERKIN-

inal incubation at 72 c for 7 minutes). As the result, a specific band was detected in 3 of the 90 pools used. One of specific band was detected in 5 sub-pools. One of these 5 a specific band was detected in 3 sub-pools. One of these 3 pools was further divided into sub-pools, each containing 30 clones, and plasmid DNA was purified from 90 sub-pools to carry out PCR in the same manner. As the result, a specific minute and synthesis at 72 c for 1 minute, followed by the pools was divided into sub-pools, each containing 250 clones, and plasmid DNA was extracted from 90 sub-pools. When these band was detected in 3 sub-pools. One of the candidate pools was cultured on the aforementioned LB plate containing 50 µg/ml of ampicillin and each of the thus formed 90 colonies these 3 pools was divided into sub-pools, each containing about 5,000 clones, and plasmid DNA was purified from 90 sub-pools to carry out PCR in the same manner. As the result, extracted samples were subjected to PCR in the same manner, was subjected to plasmid DNA extraction and PCR in the same manner. As the result, a clone HL34 was finally obtained.

<Example 16>

Sequencing of human TPO cDNA

solution and then 6 ml of 3 M potassium/5 M acetate solution pasically in accordance with the procedure described in Molecular Cloning (Sambrook et al., Cold Spring Harbor Laboratory Press, 1989). The clone HL34 was cultured overnight in 50 ml of the LB medium containing 50 µg/ml of solution. To this were added 8 ml of 0.2 N NaOH/1% SDS to suspend the cells thoroughly. After centrifugation of the shenol/chloroform (1:1), mixed with the same volume of Purification of plasmid DNA was carried out ampicillin, and the resulting cells collected by centrifugation were suspended in 4 ml of the aforementioned TEG-lysozyme suspension, the resulting supernatant fluid was treated with

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precipitation. The resulting pellet was again dissolved in the polyethylene glycol 3,000 to their final concentrations of 0.63 this way, about 300 µg of the pEF18S-HL34 plasmid DNA was sopropanol and then centrifuged. The resulting pellet was dissolved in the TE solution and treated with RNase and then with phenol/chloroform (1:1), followed by ethanol TE solution to which were subsequently added NaCl and M and 7.5%, respectively. After centrifugation, the pellet was dissolved in the TE solution and precipitated with ethanol. obtained.

using the aforementioned Taq Dye DeoxyTM Terminater Cycle The thus purified plasmid DNA was applied to the aforementioned 373A DNA sequencer (manufactured by Applied Biosystems) to determine its complete nucleotide sequence Sequening Kit (manufactured by Applied Biosystems). The thus determined nucleotide sequence and an amino acid sequence deduced therefrom are shown in the Sequence Listing (SEQ ID NO: 4). In this case, oligonucleotides synthesized based on the nucleotide sequence of SEQ ID NO: 3 and synthetic oligonucleotides designed based on the internal sequence obtained by the sequence analysis were used in the nucleotide sequence determination as primers.

open reading frame starting at its position 25, but has no 253 amino acid residues just before the poly A tail-like As the result, it was confirmed that the plasmid clone pEF18S-HL34 comprises a cDNA fragment of 861 bp and contains highly homologous sequences with the amino acid sequences AP8 (amino acid numbers 1 to 12 in SEQ ID NO: 4) analyzed in Example 2. This DNA fragment seems to encode an termination codon and contains a poly A tail-like sequence of 76 bases on its 3'-end. Its amino acid sequence, consisting of sequence, showed 84% homology with the corresponding and TP2/TP3 (amino acid numbers 157 to 162 in SEQ ID NO: 4) portion of the rat TPO cDNA (an amino acid sequence molety

cDNA. Because of the absence of a termination codon and the encodes a portion of human cDNA that corresponds to rat TPO presence of a poly A tail-like sequence on its 3'-end, this clone did not seem to be a complete cDNA, but an artificial consequence, this clone was found to be a DNA fragment which product which resulted during the procedure of cDNA library consisting of 147 residues shown in SEQ ID NO: 2). construction.

Expression of human TPO cDNA in COS 1 cells and confirmation of TPO activity <Example 17>

pEF18S-HL34 to COS 1 cells was carried out according to the Transfection of the thus obtained plasmid clone with 10 µg of the plasmid DNA by the DEAE-dextran method which includes chloroquine treatment. The COs-1 cells were cultured for 3 to 5 days at 37°c, and then the supernatant was procedure of Example 11. That is, transfection was performed collected.

extensively dialyzed against the IMDM culture medium and evaluated by the rat CFU-MK assay system. TPO activity was detected in a dose-dependent fashion in the culture supernatant of COS 1 cells in which a plasmid pEF18S-HL34 was expressed (Fig. 8). After 4 days of culturing, many contrast, TPO activity was not found in the culture supernatant of COS 1 cells in which the plasmid pEF18S alone proliferation enhancing activity was found only in the culture supernatant of COS 1 cells in which the plasmid pEF18S-HL34 was expressed through its transfection. These results The thus obtained culture supernatant was was expressed (Fig. 8). In the M-07e assay system, M-07e cell lemonstrated that pEF18S-HL34 contains a gene which encodes megakaryocytes formed elongated cytoplasmic processes.

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<Example 18>
Expression of human TPO deletion type cDNA and confirmation of TPO activity

like sequence was deleted was prepared in order to see if a protein expressed by the deletion cDNA had a TPO activity. The deletion cDNA was prepared making use of PCR. Sequences of its 3'-side, which did not exist in the rat TPO cDNA and primers used for PCR are as follows in which a restriction enzyme recognition site is added to the 5'-end of each primer EcoRI to hTPO5 and Notl and two stop codons TAA and TGA to The clone HL34 obtained in Example 15 contained the cDNA comprising a poly A tail-like continuous sequence on In consequence, a cDNA molecule from which the poly A tailtherefore seemed to be an artificial product of the experiment. h E E E

hTPO5: 5'-TTT GAA TTC GGC CAG CCA GAC ACC CCG GCC-3' (SEQ ID NO:170) (positions 1 to 21 in SEQ ID NO: 4)

htpos: 5-TTT GCG GCC GCT CAT TAT TCG TGT ATC CTG TTC AGG TAT CC-3' (SEQ ID NO:171) (anti-sense primer corresponding to positions 757 to 780 in SEQ ID NO: 4)

System 9600 (manufactured by PERKIN-ELMER) and repeating a total of 15 cycles, each cycle consisting of denaturation at with a volume of 100 µl, making use of GeneAmpTM PCR 95 c for 1 minute, annealing at 65 c for 1 minute and Using 1 µg of plasmid DNA of the plasmid clone pEF18S-HL34 obtained in Example 16 as a template, 10 µМ each of the thus synthesized hTPO5 and hTPO3 as primers and GeneAmpTM PCR Reagent Kit with AmpliTaqTM DNA Polymerase (manufactured by Takara Shuzo Co., Ltd.), PCR was carried out

incubation at 72°c for 7 minutes. A band of about 800 bp thus obtained was digested with restriction enzymes EcoRI and pEF18S which has been treated in advance with the same restriction enzymes. From the resulting transformants, 5 clones which contained a DNA fragment of about 800 bp were selected to prepare a large quantity of plasmid DNA in accordance with the procedure described in Example 5. The whole length of the amplified region of about 800 bp of each plasmid was subjected to nucleotide sequence analysis to find its complete identity with the nucleotide sequence analyzed in Notl, purified and then sub-cloned into the expression vector synthesis at 72°C for 1 minute, followed by the Example 16 (positions 1 to 780 in SEQ ID NO: 4).

under the deposit No. FERM BP-4564, in National Institute of The plasmid clone thus obtained was named pHT1-231. The vector pHT1-231 carried by an E. coli strain DH5 has been deposited by the present inventors on February 14, 1994 Bioscience and Human Technology, Agency of Industrial Science and Technology, Ministry of International Trade and Industry,

cells was carried out according to the procedure of Example 11. That is, transfection was performed with 10 µg of the plasmid DNA by the DEAE-dextran method which includes chloroquine treatment. The COS1 cells were cultured for 3 to Transfection of the thus obtained plasmid to COS 1 5 days at 37.c, and then the supernatant was collected.

found in the culture supernatant of COS 1 cells in which the plasmid pHT1-231 was expressed (Fig. 9). On 4 days of the By contrast, TPO activity was not found in the The thus obtained culture supernatant was extensively dialyzed against the IMDM culture medium and evaluated by the rat CFU-MK assay system. TPO activity was culture, many megakaryocytes formed elongated cytoplasmic culture supernatant of COS 1 cells in which the plasmid processes.

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system, M-07e cell proliferation enhancing activity was found only in the culture supernatant of COS 1 cells in which the plasmid pHT1-231 was expressed. These results demonstrated that pHT1-231 contains a cDNA which encodes a protein having In the M-07e assay DEF18S alone was expressed (Fig. 9). a TPO activity

<Example 19>

Preparation of the 3'-end region of human TPO cDNA by PCR

The clone HL34 prepared in Example 15 contained the cDNA comprising poly (A) tail-like sequence so that it was suggested that its 3'-end region was incomplete. It was The following four kinds of 5'-side primer for PCR were herefore tried to obtain a full-length 3'-end region by PCR. TPO-H: 5'-AGC AGA ACC TCT CTA GTC CTC-3' (SEQ ID NO:39) synthesized based on the sequences determined in Example 16 positions 574 to 594 in SEQ ID NO: 4)

TPO-K: 5'-ACA CTG AAC GAG CTC CCA AAC-3' (SEQ ID NO:40) positions 595 to 615 in SEQ ID NO: 4)

TPO-N: 5'-AAC TAC TGG CTC TGG GCT TCT-3' (SEQ ID NO:41) positions 660 to 680 in SEQ ID NO: 4)

1TPO-0: 5'-AGG GAT TCA GAG CCA AGA TTC-3' (SEQ ID NO:42) positions 692 to 712 in SEQ ID NO: 4)

The following 3'-side primers containing mixed nucleotides at the four bases of 3'-end were synthesized in portion. As anchor primer, without the mixed nucleotide, was order to amplify the cDNA from the beginning of the poly (A) also synthesized.

(SEQ ID NO:43) TPO3mk: 5-TAG CGG CCG C(T)17G GGG-3* (SEQ ID NO:44)

(SEQ ID NO:45) CTT-3

(SEQ ID NO:46) (SEQ ID NO:47)

hTPO3anchor: 5'-TAG CGG CCG C(T)11-3'

Jsing 1/10 volume of the synthesized cDNA solution as a emplate, 20 μM of the hTPO-H primer and 10μM of the 10 cycles, each cycle consisting of heat denaturation at 96°C for 1 minute, annealing at 48°c for 1 minute and synthesis at 3NA (manufactured by Clontech), such as in Example 14, but using 0.5 µg of oligo dT primer (it is included in TPO3mix primer and GeneAmpTM PCR Reagent Kit with AmpliTaqTM DNA Polymerase (manufactured by Takara Shuzo Co., Ltd.), PCR was carried out with volume of 50 µl, making ELMER) and heating at 96 c for 2 minutes, repeating a total of 72.c for 1 minute, followed by the final incubation at 72.c for The first strand of cDNA was synthesized from 1 TimeSaverTMcDNA Synthesis Kit manufactured by Pharmacia). use of GeneAmpTM PCR System 9600 (manufactured by PERKINug of the commercial normal human liver-derived poly (A)+ 7 minutes.

The second PCR was carried out by using 1/10 LM of the hTPO-K primer and 10µM of the hTPO3mix primer with volume of 50 µl, heating at 96 c for 2 minutes, repeating a total of 10 cycles, each cycle consisting of heat denaturation at 96.c for 1 minute, annealing at 63.c for 1 minute and volume of the first PCR resulting solution as a template, 20 synthesis at 72 c for 1 minute, followed by the final incubation at 72°c for 7 minutes.

The third PCR was carried out by using 1/10 volume of the second PCR resulting solution as a template, 20 a total of 10 cycles, each cycle consisting of heat denaturation at 96 c for 1 minute, annealing at 63 c for 1 minute and synthesis at 72.c for 1 minute, followed by the final uM of the hTPO-N primer and 10µM of the hTPO3mix primer with volume of 50 µl, heating at 96 c for 2 minutes, repeating ncubation at 72.c for 7 minutes.

The fourth PCR was carried out by using 1/10 volume of the third PCR resulting solution as a template, 20

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coincides with the nucleotide sequence of the positions 692 to 785 in SEQ ID NO: 4.

As the result of sequence analysis of the cDNA fragment in Example 17 and PCR-amplified fragment in the present Example, it is reasonably estimated that human TPO protein consists of 353 amino acids containing 21 amino-acid signal sequence, as shown in SEO ID NO: 6.

Example 20>

Reconstruction of human normal liver-derived cDNA library

Since the clone HL34 obtained in Example 15 of its open reading frame without termination codons and contained the poly A tail-like sequence directly on its 3'-end seemed therefore to be an artificial product of the cDNA synthesis; a cDNA library was reconstructed using 5 µg of a commercial normal human liver-derived poly (A)+ RNA preparation (manufactured by Clontech). Synthesis of cDNA was subjected to heat denaturation and then added to 20 µl of a reaction solution containing a Notl sequence-included oligo off as a primer attached to the kit (50 mM Tris-HCI, pH 8.3, 75 was carried out using SuperScriptTM Lambda System for cDNA Synthesis and λ Cloning Kit and SuperScript TM II RNase Hboth manufactured by LIFE TECHNOLOGIES). The poly (A)+ RNA mM KCI, 3 mM MgCl2, 1 mM DTT, 1 mM dNTP mix, 200 U SuperScriptTM II RNase H⁻), followed by 60 minutes of incubation at 37.c. After synthesis of the second strand of solution containing 25 mM Tris-HCl, pH 8.3, 100 mM KCl, 5 mM MgCl₂, 250 μM dNTP mix, 5 mM DTT, 40 U of *E. coli* DNA polymerase I, 2 U of E. coli RNase H and 10 U of E. coli DNA igase), 10 U of T4 DNA polymerase was added and the cDNA (2 hours of incubation at 16·c in 150 μl of a reaction eaction solution was heated at 65 c for 10 minutes and esulting mixture was incubated at 16 c for 5 minutes.

μM of the hTPO-O primer and 10 μM of the hTPO3mix primer with volume of 50 μl, heat at 96 c for 2 minutes, repeating a total of 10 cycles, each cycle consisting of heat denaturation at 96 c for 1 minute, annealing at 63 c for 1 minute and synthesis at 72 c for 1 minute, followed by a final incubation at 72 c for 7 minutes.

The fifth PCR was carried out by using 1/10 volume of the fourth PCR resulting solution as a template, 20 µM of the hTPO-O primer and 20µM of the hTPO3anchor primer with volume of 50 µl, heating at 96 °c for 2 minutes, repeating a total of 10 cycles, each cycle consisting of heating denaturation at 96 °c for 1 minute, annealing at 58 °c for 1 minute and synthesis at 72 °c for 1 minute, followed by the final incubation at 72 °c for 7 minutes.

The thus obtained reaction solution was subjected to 2% agarose gel (manufactured by FMC BioProducts) electrophoresis to isolate a DNA fragment of about 600 bp as the main product of the PCR which was subsequently purified making use of the aforementioned Prep-A-Gene DNA purification kit (manufactured by Bio-Rad Laboratories, Inc.). Nucleotide sequence of the thus purified DNA fragment was directly determined by a 373A DNA sequencer (manufactured by Applied Biosystems) making use of the aforementioned Taq Dye DeoxyTM Terminater Cycle Sequening Kit (manufactured by Applied Biosystems). The thus determined nucleotide sequence and an amino acid sequence deduced therefrom are shown in the Sequence Listing (SEQ ID NO: 5).

This DNA fragment has a nucleotide sequence coding for 130 amino acids starting with the primer hTPO-O, followed by sequence of more than 180 nucleotides at 3'-end (the nucleotides after 577 position could not determined). The 30 amino acid sequence starting with glycine coincides with the amino acid sequence of the positions 203 to 232 in SEQ ID NO: 4. The nucleotide sequence of positions 1 to 94 also

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(manufactured by Takara Shuzo Co., Ltd.), PCR was performed in 100 $\,\mu$ volume by GeneAmp $^{T\,M}$ PCR System 9600 carried out using 2 µg of the plasmid DNA prepared from each pool as a template and 1 µM each of the synthesized GeneAmpTM PCR Reagent Kit with AmpliTaqTM DNA Polymerase manufactured by PERKIN-ELMER) (a total of 35 cycles, each size was amplified when the plasmid DNA prepared from the each containing 15,000 transformants, and these subpools oligonucleotides (hTPO-I and hTPO-KU) as primers. Using minute, followed by the final incubation at 72.c for 7 minutes). As the result, a DNA fragment having the expected were cultured overnight in 1 ml of LB medium containing 50 using an automatic plasmid isolation apparatus PI-100. The hus extracted DNA was dissolved in TE solution, and 5% of the cycle consisting of denaturation at 95°c for 1 minute, pool # 3 was used. The # 3 pool was then divided into subpools, resulting solutions were used as templates to carry out PCR using the same primers and under the same conditions ,000 clones, and plasmid DNA was extracted and PCR was annealing at 59 c for 1 minute and synthesis at 72 c for 1 ug/ml of ampicillin, followed by extraction of plasmid DNA described above. As the result, amplification of DNA having he expected size was found in 6 of the 90 pools. When one of hese positive pools was divided into subpools, each containing parried out in the same manner as described above, DNA amplification was not observed. The density of bands on electrophoresis gels of DNA fragments amplified by a series of CR became thinner as the subpool was further subdivided, which seemed to be caused by a low recovery of the plasmid JNA due to poor growth of the clone of interest. Therefore, another screening was carried out by ybridization using the original # 3 pool.

The # 3 pool was spread on 100 LB agar plates of 15 cm in diameter in such an inoculum size that 4,100 colonies

he removal of low molecular weight DNA, attached to extracted with the same volume of phenol/chloroform, and cDNA molecules having a length of less than 400 bp were emoved using SizeSepTM 400 spun column (a spun column for 100 spun column to remove low molecular weight DNA. The After the addition of EcoRI Adaptor (attached to Directional sample was digested with Notl and again applied to SizeSepTM thus synthesized 1.3 µg of the double-stranded cDNA having an pEF18S which had been previously digested with EcoRI and TimeSaverTM cDNA Synthesis Kit manufactured by Pharmacia). Sloning Toolbox manufactured by Pharmacia), the thus treated EcoRI recognition sequence on its 5'-end and a Notl recognition sequence on its 3'-end was ligated with the expression vector Notl, and then transformed into 9.2 ml of Competent High E. coli DH5 (manufactured by Toyobo Co., Ltd.) The thus obtained numan liver cDNA library (hTPO-F1) contained 1.0 x 10⁶ ransformants.

<Example 21>

Screening of TPO cDNA clone from human liver cDNA library hTPO-F1

Human TPO cDNA-corresponding primers for PCR were synthesized based on the Sequence ID Nos. 3 and 6 shown in the SEQUENCE LISTING. Sequences of the thus synthesized orimers are as follows.

1TPO-I: 5'-TTG TGA CCT CCG AGT CCT CAG-3' (SEQ ID NO:48) positions 60 to 80 in SEQ ID NO: 3)

hTPO-KU: 5'-AGG ATG GGT TGG GGA AGG AGA-3' (SEQ ID NO:49) (anti-sense primer corresponding to the sequence of positions 301 to 921 in SEQ ID NO: 6)

transformants) constructed in Example 20 was divided into 3 pools (pool # 1 - 3) and the pools were frozen. PCR was The human liver cDNA library hTPO-F1 (1.0 x 106

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on each LB agar plate. After preparing a replica plate from each of the thus inoculated plates, one of the duplicate plates was cultured at 37°c for 6 hours to recover colonies When PCR of these DNA samples was carried out in the same manner as described above, amplification of a band equivalent o the expected size was observed in one of 100 subpools. Two and subsequent 1 hour of baking at 80°c in a vacuum oven. The supplemented with 1% SDS. Prehybridization of the thus grown on the plate and to extract the plasmid DNA samples. eplica filters were prepared from the plate of this subpool using BIODYNETM A TRANSFER MEMBRANE (manufactured by PALL). Denaturation of the filters were performed by soaking hem in 10% SDS for 10 minutes, in 0.5 N NaOH/1.5 M NaCl for 10 minutes and in 0.5 M Tris-HCI (pH 8.0)/1.5 M NaCI for 10 minutes in that order, followed by 30 minutes of air-drying thus baked filters were washed with 6 x SSC (prepared by and 88.2 g of Na citrate dissolved in 1 liter water, pH 7.0) washed filters were carried out by incubating it at 42.c for 30 ninutes with shaking in 30 ml of a reaction solution consisting of 50% formamide, 5 x SSC, 5 x Denhardt's solution prepared from 50 x Denhardt's solution containing 5 g of diluting 20 x SSC stock solution consisting of 175.3 g of NaCl albumin fraction V in 500 ml water), 1% SDS and 20 µg/ml of salmon sperm DNA. After prehybridization, the reaction and then incubated with shaking at 42 c for 20 hours. The ourifying a portion of SEQ ID NO: 4 ranging from its 5'-end to icoll, 5 g of polyvinylpyrrolidone and 5 g of bovine serum solution was exchanged with 30 ml of a hybridization solution naving the same composition, mixed with a probe which had ragment of the plasmid pEF18S-HL34, which was obtained by he 458 position base and by labeling the purified portion by been labeled with $[\alpha-3^2P]$ dCTP (manufactured by Amersham) abeled probe used in this experiment was the EcoRI/BamHI

random primer technique with Megaprime DNA Labelling System (a kit manufactured by Amersham based on the method disclosed in Anal. Biochem., 132, 6 - 13, 1983). The thus reated filters were washed in 2 x SSC/0.1% SDS solution at 12 c for 30 minutes and then in 0.2 x SSC/0.1% SDS solution at 42.c for 30 minutes. The filters were then subjected to 16 hours of autoradiography at -70 c using an intensifying screen were collected from the original plate and inoculated again onto a 10-cm LB agar plate. A total of 50 colonies grown on the plate were separately cultured, and their DNA samples As the result, a single signal considered to be positive was bserved. Colonies approximately corresponding to this signal were subjected to PCR using the aforementioned primers hTPO-I and hTPO-KU under the same conditions as described above. As the result, amplification of a band equivalent to the expected size was found in only one clone, which was and an X-OMATTM AR5 film (manufactured by Eastman Kodak). designated pHTF1.

Example 22>

Determination of nucleotide sequence of human TPO CDNA clone pHTE1

Purification of plasmid DNA was carried out basically according to the procedure described in *Molecular Cloning* (Sambrook *et al.*, Cold Spring Harbor Laboratory Press, 1989). The clone pHTF1 was cultured overnight in 50 ml of the LB medium containing 50 µg/ml of ampicillin. The resulting cells were collected by centrifugation and suspended in 4 ml of the aforementioned TEG-lysozyme solution. To this were added 8 ml of 0.2 N NaOH/1% SDS solution and then 6 ml of 3 M potassium/5 M acetate solution to suspend the cells thoroughly. After centrifugation of the suspension, the resulting supernatant was treated with phenol/chloroform (1:1), mixed with the same volume of isopropanol and then

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centrifuged. The resulting pellet was dissolved in TE solution and treated with RNase and then with phenol/chloroform (1:1), followed by ethanol precipitation. The resulting pellet was again dissolved in TE solution to which were subsequently added NaCl and polyethylene glycol 3,000 to their final concentrations of 0.63 M and 7.5%, respectively. After sentritugation, the pellet was dissolved in TE solution and precipitated with ethanol. In this way, about 300 µg of the plasmid DNA pHTF1 was obtained.

The thus purified plasmid DNA was applied to the aforementioned 373A DNA sequencer (manufactured by Applied Biosystems) to determine its complete nucleotide sequence using the aforementioned Taq Dye DeoxyTM Terminater Cycle Sequening Kit (manufactured by Applied Biosystems). The thus determined nucleotide sequence and the deduced amino acid sequence are shown in the Sequence Listing (SEQ ID NO: 7). Oligonucleotides synthesized based on the nucleotide sequence of SEQ ID NO: 6 and on the internal sequence obtained by their sequence. reaction analysis were used in the nucleotide sequence determination as primers.

As the result, it was confirmed that the clone pHTF1 contains a cDNA fragment of 1,721 bp and has high homology with the amino acid sequences AP8 (amino acid numbers 1 to 12 in SEQ ID NO: 7) and TP2/TP3 (amino acid numbers 157 to 162 in SEQ ID NO: 7) analyzed in Example 2. This DNA fragment seems to have a 5' noncoding region of 101 bases, an open reading frame consisting of 353 amino acid residues, starting at a methionine residue encoded at the 102 to 104 nucleotide positions and ending at a glycine residue encoded at the 1,158 to 1,160 nucleotide positions, the subsequent termination codon (TAA), a 3' noncoding region of 531 bases and a poly A tail sequence of 30 bases. The amino acid sequence of a protein considered to be encoded by the open reading frame completely coincided with the predicted amino

nucleotide sequence was different from SEQ ID NO: 6 at 3 sequence of pHTF1 has a larger size than the putative cDNA sequence shown in SEQ ID NO: 6, and contains 77 more additional bases at the 5' side and 347 more additional bases at the 3' side in front of its poly A tail sequence. Also, the positions. That is, A (position 84), A (position 740) and G icid sequence of human TPO shown in SEQ ID NO: 6. The cDNA position 1,198) in SEQ ID NO: 7 were C, T and A in SEQ ID NO: 6, respectively. Only the mutation at position 740 was included in the protein-encoding region, but this mutation did not cause amino acid exchange because both bases A and T were the third base of threonine codons. Although the cause of these base substitutions was not clear at that time, analysis of the plasmid clone pHTF1 confirmed that human TPO protein comprised 353 amino acid residues having a 21 amino acid signal sequence. Molecular weight of the mature protein excluding the signal sequence was estimated to be 35,466.

The vector pHTF1 carried by an *E. coli* strain DH5 nas been deposited by the present inventors on March 24, 1994 under the accession No. FERM BP-4617, at the National Institute of Bioscience and Human-Technology, Agency of Industrial Science and Technology, Ministry of International Irade and Industry, Japan.

<Example 23>

Expression of human TPO cDNA clone pHTF1 in COS 1 cells and confirmation of TPO activity

Transfection to COS 1 cells of the thus obtained plasmid clone pHTF1 was carried out according to the procedure of Example 11. That is, transfection was performed with 10 µg of the plasmid DNA by the DEAE-dextran method which includes chloroquine treatment. The transfected COS1

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The culture supernatant was extensively dialyzed against the IMDM culture medium and evaluated by the rat CFU-MK assay system. TPO activity was detected in a dose-dependent fashion in the supernatant of COS1 cells in which the plasmid pHTF1 was expressed (Fig. 10a). By contrast, TPO activity was not found in the culture supernatant of COS1 cells in which the plasmid pEF18S alone was expressed (Fig. 10a). Similar results were obtained in the M-07e assay system. The supernatant of COS1 cells, in which the plasmid pHTF1 was expressed, significantly augmented M-07e cell proliferation in a dose-dependent manner (Fig. 10b). These results demonstrated that pHTF1 contains a gene which encodes a protein having TPO activity.

Example 24>

Cloning of human TPO chromosomal DNA

carried out using human TPO cDNA as a probe. A genomic ibrary used in the cloning was a gift from Prof. T. Yamamoto at the Gene Research Center, Tohoku University (the library 1994, constructed by partially digesting human chromosomal DNA with a restriction enzyme Sau3AI and ligating the partial digest to the BamHI site of a phage vector Lambda EMBL3 manufactured by Stratagene). Screening from this library using human TPO cDNA as a probe was carried out basically according to the procedure described in Molecular Cloning Using E. coli LE392 as a host, the library was inoculated onto a 15-cm plate containing NZYM (10 g of NZ amine, 5 g of NaCl, 5 g of Bacto Yeast Extract, 2 g of MgSO4.7H2O and 15 g of agar in 1 liter water, pH 7.0) in such an inoculum size that one plate . Cloning of human TPO chromosomal DNA was reported by Sakai et al. in J. Biol. Chem., 269, 2173 - 2182, Sambrook et al., Cold Spring Harbor Laboratory Press, 1989)

HCI (pH 8.0)/1.5 M NaCl for 10 minutes, followed by 30 minutes of air-drying and subsequent 1.5 hours of baking at 80°C in a reaction solution consisting of 50% formamide, 5 x SSC, 5 x by PCR and purified, and the purified fragment was labeled with 32p making use of Random Primer DNA Labelling Kit (a contained 30,000 phage particles. Two replica filters were prepared from each of the thus prepared 18 plates using Denaturation of the filters was performed by soaking them in D.5 N NaOH/1.5 M NaCl for 10 minutes and then in 0.5 M Trisvacuum oven. Prehybridization was carried out by incubating the thus treated filters at 42 c for 1 hour in 500 ml of a Denhardt's solution, 1% SDS and 20 µg/ml of salmon sperm DNA. For use as a probe, a human TPO cDNA fragment (base position numbers 178 to 1,025 in SEQ ID NO: 7) was amplified DNA labeling kit manufactured by Takara Shuzo based on the random primer method disclosed in Anal. Biochem., 132, 6 - 13, BIODYNETM A TRANSFER MEMBRANE (manufactured by PALL). htpo-1; 5'-ttg tga cct ccg agt cct cag-3' (SEQ ID NO:50) 1983). Sequences of primers used in this PCR are as follows. (positions 178 to 198 in SEQ ID NO: 7) (anti-sense primer corresponding to the sequence of positions 1,005 to 1,025 in SEQ ID NO: 7)

Using the isotope-labeled probe, hybridization was carried out at 42° c for 20 hours in 500 ml of a reaction solution having the same composition of the prehybridization solution. The resulting filters were washed 3 times in 2 x SSC/0.1% SDS solution at room temperature for 5 minutes, and

TPO-N: 5'-AGG GAA GAG CGT ATA CTG TCC-3' (SEQ ID NO:51)

then once in 0.1 x SSC/0.1% SDS solution at 68°C for 1 hour. The filters were then subjected to 16 hours of autoradiography at -70°C using an intensifying screen and an X-OMATTM AR5 film (manufactured by Eastman Kodak). As the result, 13 positive signals were obtained. Plaques approximately corresponding to each of the positive signals were collected



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from the original plates and inoculated again onto 15-cm NZYM plates in such an inoculum size that 1,000 plaques were formed on each plate. Two replica filters were prepared from each of the resulting plates to carry out hybridization under the same conditions described above. As the result, positive signals were detected on all filters of the 13 groups. A single plaque was recovered from each of the resulting plates to prepare phage DNA by the plate lysate method described in Molecular Cloning. Phage DNA samples thus prepared from the 13 clones were checked for the presence of the cDNA coding region, by PCR using primers having the following sequences. hTPO-L: 5'-GGC CAG CCA GAC ACC CCG GCC-3' (SEQ ID NO:52) (positions 1 to 21 in SEQ ID NO: 6)

hTPO-F: 5-ATG GGA GTC ACG AAG CAG TTT-3' (SEQ ID NO:53) (anti-sense primer corresponding to the sequence of positions 127 to 147 in SEQ ID NO: 6)

htpo-p: 5'-tgc Gtt tcc tga tgc ttg tag-3' (Seq Id NO:54) positions 503 to 523 in Seq Id NO: 6)

hTPO-V: 5'-AAC CTT ACC CTT CCT GAG ACA-3' (SEQ ID NO:55) (anti-sense primer corresponding to the sequence of positions 1,070 to 1,090 in SEQ ID NO: 6)

When PCR was carried out using these primer combinations, 5 of the 13 clones seemed to contain entire amino acid coding region predicted from the cDNA. Chromosomal DNA molecules contained in these 5 clones had a similar length of about 20 kb and showed almost the same pattern when a preliminary restriction enzyme analysis was performed. Therefore, one of these clones (clone λHGT1) was selected and analyzed by Southern blotting. That is, 1 μg of DNA of the clone λHGT1 was digested completely with a restriction enzyme *EcoR*I or *Hind*III and applyeded to 0.8% agarose gel electrophoresis, and bands on the gel were transferred on BIODYNETM A TRANSFER MEMBRANE (manufactured by PALL). The resulting filter was air-dried for

30 minutes and subjected to 2 hours of baking at 80°C in a DNA. For use as a probe, a human TPO cDNA fragment (base by PCR and purified, and the purified fragment was labeled nl of a reaction solution having the same composition of the prehybridization solution. The resulting filter was washed 3 limes in 2 x SSC/0.1% SDS solution at room temperature for 5 X-OMATTM AR5 film (manufactured by Eastman Kodak). As the vacuum oven. Prehybridization was performed by incubating the thus treated filter at 42°C for 1 hour in 50 ml of a reaction solution consisting of 50% formamide, 5 x SSC, 5 x Denhardt's solution, 1% SDS and 20 µg/ml of salmon sperm position numbers 178 to 1,025 in SEQ ID NO: 7) was amplified with 32P using Random Primer DNA Labelling Kit (manufactured by Takara Shuzo). Using the isotope-labeled probe, hybridization was carried out at 42°c for 20 hours in 50 autoradiography at -70 c using an intensifying screen and an esult, a single band of about 10 kb was observed in the case of the Hindlil digestion. Therefore, 10 µg of DNA of the clone AHGT1 was digested with HindIII and subjected to 0.8% agarose gel electrophoresis, and the 10 kb band was excised from the ligested with HindIII. In this case, Competent-High E. coli DH5 minutes, and then once in 0.1 x SSC/0.1% SDS solution at 68 c or 1 hour. The filter was then subjected to 16 hours of jeł, purified using Prep-A-Gene DNA Purification Kit (manufactured by Bio-Rad) and subcloned into a cloning vector JUC13 (manufactured by Pharmacia) which had been previously nanufactured by TOYOBO was used as the host strain.

Of the thus obtained clones, a clone containing the 10 kb HindIII fragment was selected and designeted pHGT1.

A restriction map of phage clone λHGT1 is shown ig.11.

<Example 25>

Determination of nucleotide sequence

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of human TPO chromosomal clone pHGT1

plasmid DNA were carried out basically according to the method described in Molecular Cloning (Sambrook et al., Cold Spring Harbor Laboratory Press, 1989). The clone pHGT1 was ug/ml of ampicillin. The resulting cells were collected by same volume of isopropanol and then centrifuged. The cultured overnight in 50 ml of the LB medium containing 50 FEG-lysozyme solution. To this were added 8 ml of 0.2 N VaOH/1% SDS solution and then 6 ml of 3 M potassium/5 M acetate solution to suspend the cells thoroughly. After was treated with phenol/chloroform (1:1), mixed with the resulting pellet was dissolved in the TE solution and treated dissolved in the TE solution and precipitated with ethanol. In centrifugation of the suspension, the resulting supernatant in the TE solution to which were subsequently added NaCl and M and 7.5%, respectively. After centrifugation, the pellet was his way, about 300 µg of the plasmid DNA pHGT1 was centrifugation and suspended in 4 ml of the aforementioned with RNase and then with phenol/chloroform (1:1), followed by ethanol precipitation. The resulting pellet was again dissolved polyethylene glycol 3,000 to their final concentrations of 0.63 Culturing of the clone pHGT1 and purification

Using the aforementioned Taq Dye Deoxy^{T M} Biosystems), the thus purified plasmid DNA was applied to the sequence. The thus determined nucleotide sequence and the ferminater Cycle Sequening Kit (manufactured by Applied aforementioned 373A DNA sequencer (manufactured by Applied Biosystems) to determine its nucleotide sequence around the protein-encoding region predicted from the cDNA nucleotide deduced amino acid sequence are shown in the Sequence Listing SEQ ID NO: 8). In this case, oligonucleotides used in Example 22 for the nucleotide sequence analysis of cDNA and synthetic

obtained by their sequence reaction analysis were used as oligonucleotides designed based on the internal sequence primers in the nucleotide sequence determination.

As the result, chromosomal DNA carried by the plasmid clone pHGT1 was found to contain an entire coding and nucleotide sequence of the coding region coincided completely with that of SEQ ID NO: 6. In addition, a region ntrons having lengths of 231 bp, 286 bp, 1,932 bp and 236 bp nucleotide sequence was different from the cDNA nucleotide egion of the amino acid sequence deduced from SEQ ID NO: 6, corresponding to the amino acid-encoding exon contained 4 in that order counting from the 5' side (Fig. 11). Also, the sequence of SEQ ID NO: 7 at 3 positions which were the same as the positions described in Example 22 (different positions between Sequence ID Nos. 6 and 7). That is, A (position 84), A position 740) and G (position 1,198) in SEQ ID NO: 7 was respectively C, T and A in SEQ ID NO: 8. Thus, it was revealed hat the nucleotide sequence of the human TPO cDNA clone oHTF1 obtained in Example 21 is different from the nucleotide sequence of the human chromosomal DNA clone pHGT1 at 3 positions. Therefore, in order to analyze if these mutations in he cDNA clone pHTF1 sequence reflect the chromosomal DNA sequence, nucleotide sequences of other 4 clones among the 5 chromosomal DNA clones independently selected by the screening were determined. The sequence analysis was carried out by a direct nucleotide sequence determination method using a phage DNA sample prepared according to the plate using the sequence primers used in Example 22, which have been synthesized based on such sequence portions that the iforementioned 3 mutation positions in the nucleotide ysate method described in Molecular Cloning. Sequencing of each clone was performed by applying it to the aforementioned equence could be analyzed, and the aforementioned Taq Dye 373A DNA sequencer (manufactured by Applied Biosystems),

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of all 4 clones were identical to that of SEQ ID NO: 6. The Applied Biosystems). It was found that nucleotide sequences positions 84 and 740 corresponding to SEQ ID NO: 7 were substituted by C and T, respectively, in these 4 clones. But the DeoxyTM Terminater Cycle Sequening Kit (manufactured by position 1,198 was G in 2 clones and A in other 2 clones. In other words, it was revealed that there are two types of nucleotide sequences inherent to the original chromosomal DNA. At present, it is not clear if such differences in the nucleotide sequence are derived from homologous chromosome or from plural genes. In addition, it is suggested that the differences in the nucleotide sequence may be due to a racial difference, because the poly (A)+ RNA purchased from Clontech is a Caucasian origin while the chromosomal DNA is a Japanese

The vector pHGT1 carried by an E. coli strain DH5 has been deposited by the present inventors on March 24, 1994 Institute of Bioscience and Human-Technology, Agency of under the accession No. FERM BP-4616, at the National Industrial Science and Technology, Ministry of International Frade and Industry, Japan.

<Example 26>

Expression of human TPO chromosomal DNA in COS 1 cells and confirmation of TPO activity The plasmid clone pHGT1 obtained by the subcloning contained a total of 4 EcoRI recognition sequences, 3 in the insert moiety and 1 in the vector. The nucleotide sequence region was included in a DNA fragment of about 4.3 kbp which was interposed between the EcoRI recognition sequence closest to the 5' side in the insert and the EcoRI recognition sequence in the vector. Therefore, this fragment was ligated analysis revealed that the entire human TPO protein-encoding

preparing these plasmid DNA, an expression experiment was carried out. Purification of plasmid DNA was carried out with an EcoRI-treated expression vector pEF18S, and obtained 4 human TPO expression plasmid pEFHGTE#1-4 (see Fig. 11). By basically according to the procedure described in Molecular Cloning (Sambrook et al., Cold Spring Harbor Laboratory Press, 989), thereby obtaining about 250 µg of the plasmid DNA.

Transfection to COS 1 cells of the thus obtained clones pEFHGTE#1-4 was carried out according to the procedure of Example 11. That is, transfection was performed with 10 µg of the plasmid DNA by the DEAE-dextran method cells were incubated for 3 days at 37 c and then the culture which includes chloroquine treatment. The transfected COS1 supernatants were collected.

The culture supernatants were extensively dialyzed against the IMDM culture medium and evaluated by the rat CFU-MK assay system. TPO activity was detected in a dosedependent fashion in the supernatant of COS1 cells in which TPO activiy was not found in the supernatant of COS1 cells in Similar results were obtained in the M-07e assay system. The supernatant of COS1 cells, in which each of four clones (PEFHGTE#1-4) was expressed, significantly augmented M-07e cell growth in a dose-dependent fashion. Representative data each of four clones (pEFHGTE#1-4) was expressed. By contrast, Representative data with pEFHGTE#1 is shown in Fig. 12a. which the plasmid pEF18S alone was expressed. with pEFHGTE#1 is shown in Fig. 12b.

clones pEFHGTE#1-4 carry the functional chromosomal DNA of These results demonstrated that the human TPO.

<Example 27>

expression in COS 1 cells and confirmation of TPO activity Preparation of human TPO deletion type DNA and its

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encoded by pHT1-231 were examined for their ability to exert emoval of its carboxyl third. Thus, in order to further derivatives lacking 20, 40, 60 or 68 amino acids from the TPO could exhibit its biological activity even after the evaluate the biologically active portions, deletion derivative experiments were performed. In this Example, TPO deletion carboxyl-terminal end of the TPO protein (amino acids 1-231) IPO in vitro biological activity. The shortest derivative corresponding to TP2/3 of rat plasma TPO described in amino acids 1-163) still included amino acid sequences Example 2. The deletion plasmids were prepared by PCR using he DNA of the plasmid clone pHT1-231 obtained in Example 18 The results in Example 18 indicated that human as template and synthesized oligonucleotides as primers.

htpo-5: 5'-TTT GAA TTC GGC CAG CCA GAC ACC CCG GCC-3' SEQ ID NO:56) (prepared by adding an EcoRI recognition sequence to the sequence of positions 1 to 21 in SEQ ID NO:4; Sequences of primers used for PCR were as follows: his is the identical sequence described in Fig.9);

GTG GGT-3' (SEQ ID NO:57) (an antisense primer corresponding prepared by adding two termination codons TAA and TGA and a VotI recognition sequence for use in the preparation of a deletion derivative coding for positions 1 to 163 amino acid hTPO-S: 5'-TTT GCG GCC GCT CAT TAG CTG GGG ACA GCT GTG to the sequence of positions 555 to 576 in SEQ ID NO:4, esidues):

TPO4: 5-TTT GCG GCC GCT CAT TAC AGT GTG AGG ACT AGA 3AG GTT CTG-3' (SEQ ID NO:58) (an antisense primer corresponding to the sequence of positions 576 to 600 in SEQ D NO:4, prepared by adding two termination codons TAA and IGA and a Not recognition sequence for use in the preparation of a deletion derivative coding for positions 1 to 171 amino icid residues);

hTPO-30: 5'-TTT GCG GCC GCT CAT TAT CTG GCT GAG GCA GTG corresponding to the sequence of positions 636 to 660 in SEQ ID NO:4, prepared by adding two termination codons TAA and TGA and a Not recognition sequence for use in the preparation of a deletion derivative coding for positions 1 to 191 amino AAG TTT GTC-3' (SEQ ID NO:59) (an antisense primer acid residues); and

hTPO-2: 5'-TTT GCG GCC GCT CAT TAC AGA CCA GGA ATC TTG corresponding to the sequence of positions 696 to 720 in SEQ ID NO:4, prepared by adding two termination codons TAA and TGA and a Not recognition sequence for use in the preparation GCT CTG AAT-3' (SEQ ID NO:60) (an antisense primer of a deletion derivative coding for positions 1 to 211 amino acid residues).

followed by the final incubation at 72 c for 7 minutes. A band obtained in Example 18 as a template, 10 µM each of the thus synthesized hTPO-5 (for 5' side) and hTPO-2, -3, -4 and -S (for PCR was carried out with a volume of 100 µl, using ELMER) and repeating a total of 20 cycles, each cycle consisting of denaturation at 95°c for 1 minute, annealing at naving an expected size amplified by each PCR. The thus 3' side) as primers and GeneAmpTM PCR Reagent Kit with AmpliTaqTM DNA Polymerase (manufactured by Takara Shuzo), GeneAmpTM PCR System 9600 (manufactured by PERKINof interest thus obtained by each PCR was digested with estriction enzymes EcoRI and NotI and the resulting digest was subjected to 1% agarose gel (manufactured by FMC BioProducts) electrophoresis to isolate a main DNA fragment solated DNA fragment was purified using Prep-A-Gene DNA sloned into the expression vector pEF18S which had been previously treated with the same restriction enzymes. In this Using 1 µg of plasmid DNA of the clone pHT1-231 35 c for 1 minute and synthesis at 72 c for 1 minute, Jurification Kit (manufactured by Bio-Rad) and then sub-

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samples basically according to the procedure described in case, Competent-High E. coli DH5 manufactured by TOYOBO was From the resulting transformants which originated from each PCR, 4 to 5 clones containing the nsert of expected size were selected to prepare plasmid DNA Molecular Cloning (Sambrook et al., Cold Spring Harbor aboratory Press, 1989). In a series of such operations, plasmid DNA samples were obtained from a deletion derivative coding for positions 1 to 163 amino acid residues (pHT1-163 amino acid residues (pHT1-171 #1-4), a deletion derivative coding for positions 1 to 191 amino acid residues (pHT1-191 #1-4) and a deletion derivative coding for positions 1 to 211 amino acid residues (pHT1-211 #1-4). The whole length of the amplified regions of each plasmid DNA was subjected to nucleotide sequence analysis to find its complete identity #1-5), a deletion derivative coding for positions 1 to 171 with the nucleotide sequence shown in SEQ ID NO:4. as the host strain.

clones was carried out according to the procedure of Example 11. That is, transfection was performed with 10 µg of each of the plasmid DNA samples by the DEAE-dextran method which includes chloroquine treatment. The transfected COS1 cells were cultured for 3 days and then the culture supernatants Transfection to COS 1 cells of the thus obtained were recovered.

The culture supernatants were extensively dialyzed 191#1 and pHT1-171#2 is shown in Fig. 13a and with pHT1-163#2 in Fig. 13b. Similar results were obtained in the M-07e assay system. The supernatant of COS1 cells, in which pHT1against the IMDM culture medium and evaluated by the rat CFU-MK assay system. TPO activity was detected in a dosedependent fashion in the supernatant of COS1 cells in which pHT1-211, pHT1-191, pHT1-171 or pHT1-163 each was expressed. Representative data with pHT1-211#1, pHT1-211, pHT1-191, pHT1-171 or pHT1-163 each was expressed,

proliferation. Representative data with pHT1-211#1, pHT1-191#1, pHT1ee Ce 171#2 or pHT1-163#2 is shown in Fig. 14. augmented M-07e significantly

These results showed that human TPO still retains erminal half beyond Ser (position 163) is deleted, and strongly suggested that the biologically active portions of human TPO localize within the amino-terminal half ending at ts in vitro biological activity even after its carboxyl-Ser (position 163).

<Example 28>

Preparation of C-terminal deletion type human TPO and its expression and activity in COS 1 cells

amino acids were further deleted from the deletion derivatives In order to analyze the region necessary for the human TPO protein that exhibits its activity, the C-terminal obtained in Example 27 and the TPO activity expressed in COS 1 cells was examined. Using the plasmid clone pEF18S-HL34 obtained in Example 16, expression plasmids were prepared by deleting nucleotide sequences coding for C-terminal amino acid residues starting from the 163 position serine. Construction of these deletion plasmids was effected making use of PCR. Sequences of primers prepared for use in PCR are as follows:

CCG GCC-3' (SEQ ID NO:61) (prepared by adding an EcoRI recognition sequence to the sequence of positions 1 to 21 in htpo-5: 5'-TTT GAA TTC GGC CAG CCA GAC ACC SEQ ID NO: 4; the same sequence shown in Example 18);

hTPO-150: 5'-TTT GCG GCC GCT CAT TAG AGG GTG GAC CCT CCT ACA AGC AT-3' (SEQ ID NO:62) (an antisense primer corresponding to the sequence of positions 514 to 537 in SEQ ID NO:4; prepared by adding two termination codons TAA and TGA and a Notl recognition sequence for use in the - 150 -

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preparation of a deletion mutant coding for positions 1 to 150 amino acid residues);

htpo-151: 5'-TTT GCG GCC GCT CAT TAG CAG AGG GTG GAC CCT CCT ACA A-3' (SEQ ID NO:63) (an antisense primer corresponding to the sequence of positions 518 to 540 in SEQ ID NO:4; prepared by adding two termination codons TAA and TGA and a Not recognition sequence for use in the preparation of a deletion derivative coding for positions 1 to 151 amino acid residues);

hTPO-153: 5'-TTT GCG GCC GCT CAT TAC CTG ACG corresponding to the sequence of positions 526 to 546 in SEQ IGA and a Noti recognition sequence for use in the preparation CAG AGG GTG GAC CC-3' (SEQ ID NO:64) (an antisense primer D NO:4; prepared by adding two termination codons TAA and of a deletion derivative coding for positions 1 to 153 amino acid residues);

corresponding to the sequence of positions 529 to 549 in SEQ D NO:4, prepared by adding two termination codons TAA and hTPO-154: 5'-TTT GCG GCC GCT CAT TAC CGC CTG ACG CAG AGG GTG GA-3' (SEQ ID NO:65) (an antisense primer IGA and a Not recognition sequence for use in the preparation of a deletion derivative coding for positions 1 to 154 amino acid residues);

corresponding to the sequence of positions 532 to 552 in SEQ ID NO:4, prepared by adding two termination codons TAA and IGA and a Not recognition sequence for use in the preparation of a deletion derivative coding for positions 1 to 155 amino htpo-155; 5-TTT GCG GCC GCT CAT TAG GCC CGC CTG ACG CAG AGG GT-3' (SEQ ID NO:66) (an antisense primer acid residues);

corresponding to the sequence of positions 535 to 555 in SEQ ID NO:4, prepared by adding two termination codons TAA and htpo-156: 5'-TTT GCG GCC GCT CAT TAT GGG GCC CGC CTG ACG CAG AG-3' (SEQ ID NO:67) (an antisense primer

IGA and a Not recognition sequence for use in the preparation of a deletion derivative coding for positions 1 to 156 amino acid residues); and

htpo-157: 5'-TTT GCG GCC GCT CAT TAG GGT GGG corresponding to the sequence of positions 538 to 558 in SEQ ID NO.4, prepared by adding two termination codons TAA and IGA and a Notl recognition sequence for use in the preparation of a deletion derivative coding for positions 1 to 157 amino GCC CGC CTG ACG CA-3' (SEQ ID NO:68) (an antisense primer acid residues).

(hTPO-5 for 5' side and hTPO-150, -151, -153, -154, -155, the clone pEF18S-HL34 obtained in Example 16 as a template and 10 µM each of the thus synthesized oligonucleotides 56 and -157 for 3' side) as primers. Using GeneAmpTM PCR (manufactured by FMC BioProducts) electrophoresis to isolate Reagent Kit with AmpliTaqTM DNA Polymerase (manufactured volume making use of GeneAmpTM PCR System 9600 manufactured by PERKIN-ELMER) and repeating a total of 20 minute, annealing at 66°C for 1 minute and synthesis at 72°C minutes. A band of interest thus obtained by each PCR was digested with restriction enzymes EcoRI and Notl and the a main DNA fragment having a size expected by each PCR. The hus isolated DNA fragment was purified using Prep-A-Gene reated in advance with the same restriction enzymes. In this case, Competent-High E. coli DH5 manufactured by TOYOBO was From the resulting transformants PCR was carried out using 1 µg of plasmid DNA of by Takara Shuzo), the PCR reaction was performed in 100 μl for 1 minute, followed by the final incubation at 72°c for 7 resulting digest was subjected to 1% agarose gel DNA Purification Kit (manufactured by Bio-Rad) and then subcloned into the expression vector pEF18S which has been which originated from each PCR experiment, 3 to 5 clones cycles, each cycle consisting of denaturation at 95°c for 1 used as the host strain.

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containing the insert of expected size were selected to prepare plasmid DNA samples basically in accordance with the procedure described in Molecular Cloning (Sambrook et al., Cold operations, plasmid DNA samples were obtained from a deletion derivative coding for positions 1 to 150 amino acld 16, 17 and 18), a deletion derivative coding for positions 1 to (pHT1-154 # 1 to 5), a deletion derivative coding for positions Spring Harbor Laboratory Press, 1989). By a series of such esidues (pHT1-150 # 21, 22 and 25), a deletion derivative coding for positions 1 to 151 amino acid residues (pHT1-151 # derivative coding for positions 1 to 154 amino acid residues derivative coding for positions 1 to 156 amino acid residues 153 amino acid residues (pHT1-153 # 1 to 5), a deletion to 155 amino acid residues (pHT1-155 # 1 to 5), a deletion (pHT1-156 # 1 to 5) and a deletion derivative coding for positions 1 to 157 amino acid residues (pHT1-157 # 1 to 5).

Of these purified plasmid DNA samples, the sequence of pHT1-150 # 21, 22 and 25 and pHT1-151 # 16, 17 and 18 were checked by 373A DNA Sequencer manufactured by Cycle Sequening Kit (Applied Biosystems), thereby confirming expected TPO cDNA sequences with no substitutions over the Applied Biosystems making use of Taq Dye DeoxyTM Terminater entire nucleotide sequences.

evaluated by the M-07e assay system. TPO activities were Transfection of GOS 1 cells with each of the thus obtained clones was carried out in accordance with the dextran method which includes chloroquine treatment, and the he IMDM culture medium described in the foregoing and detected in the culture supernatant of COS 1 cells transfected procedure of Example 11. That is, transfection was performed with 10 µg of each of the plasmid DNA samples by the DEAE-The thus obtained culture supernatants were dialyzed against with respective clones coding for C-terminal deletion culture supernatant was recovered after 3 days of the culture.

153 position amino acids, 1 to 154 position amino acids, 1 to to 157 position amino acids. All of these derivatives contain a were detected in the culture supernatant of COS 1 cells transfected with the clone coding for a C-terminal side cysteine residue at position 151. However, no TPO activities deletion derivative consisting of 1 to 150 position amino acids 155 position amino acids, 1 to 156 position amino acids or 1 derivatives, consisting of 1 to 151 position amino acids, 1 whose cysteine residue at the 151 position was also deleted.

Preparation of N-terminal deletion type human TPO and its expression and activity in COS 1 cells <Example 29>

terminal side amino acids of the deletion derivatives obtained in Example 28 were further deleted and expression of TPO examined. Using the plasmid clone pEF18S-HL34 obtained in activity of the thus prepared deletion derivativess was Example 16 and the plasmid clone pHT1-163 obtained in nucleotide sequences coding for N-terminal amino acid residues after the signal sequence. Construction of these In order to analyze the region necessary for the TPO protein that exhibits its biological activity, the N-Example 27, expression plasmids were prepared by deleting deletion plasmids were effected making use of PCR. Sequences of primers prepared for use in PCR are as follows:

hTPO-5: 5'-TTT GAA TTC GGC CAG CCA GAC ACC recognition sequence to the sequence of positions 1 to 21 in CCG GCC-3' (SEQ ID NO:69) (prepared by adding an *Eco*RI SEQ ID NO:4; the same sequence shown in Example 18);

hTPO3: 5'-TTT GCG GCC GCT CAT TAT TCG TGT ATC CTG TTC AGG TAT CC-3' (SEQ ID NO:70) (an antisense primer ID NO:4; prepared by adding two termination codons TAA and corresponding to the sequence of positions 757 to 780 in SEQ

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GGA CAG CGT TAG CCT TGC AGT TAG-3' (SEQ ID NO:77) (an antisense primer corresponding to the sequence of positions 64 to 87 and 109 to 132 in SEQ ID NO:4, prepared for use in the htpo-8R: 5'-CAG TTT ACT GAG GAC TCG GAG GTC preparation of a derivative in which amino acid residues of positions 1 to 7 are deleted).

in which amino (1) Preparation of a derivative (pHT13-231) acid residues of positions 1 to 12 are deleted

the clone pEF18S-HL34 obtained in Example 18 as a template and 5 µM each of the thus synthesized oligonucleotides (hTPO-13 and hTPO3 in one combination and hTPO-5 and hTPO-13R in another combination) as primers. Using GeneAmpTM PCR PCR was carried out using 1.4 µg of plasmid DNA of by Takara Shuzo), the PCR reaction was performed in 100 µl (manufactured by PERKIN-ELMER) and, after 5 minutes of annealing at 65 c for 1 minute and synthesis at 72 c for 1 (manufactured by FMC BioProducts) electrophoresis to isolate a main DNA fragment having an expected size which was Reagent Kit with AmpliTaqTM DNA Polymerase (manufactured volume making use of GeneAmpTM PCR System 9600 denaturation at 95°c, repeating a total of 30 cycles, each Each of the PCR products was subjected to 1.2% agarose gel subsequently purified using Prep-A-Gene DNA Purification Kit Thereafter, second PCR was carried out using 1 µl portion of cycle consisting of denaturation at 95°c for 1 minute, minute, followed by the final incubation at 72°c for 7 minutes. manufactured by Bio-Rad) and dissolved in 15 µl of TE buffer. each of the thus prepared solution as a template.

The second PCR was carried out using 5 µM each of GeneAmpTM PCR Reagent Kit with AmpliTaqTM DNA Polymerase manufactured by Takara Shuzo), the PCR reaction was he synthesized primers (hTPO-5 and hTPO3).

GA and a Notl recognition sequence; the same sequence synthesized for use in the preparation of a deletion derivative coding for positions 1 to 231 amino acid residues);

ID NO:4; prepared for use in the preparation of a deletion hTPO-S: 5-TTT GCG GCC GCT CAT TAG CTG GGG ACA GCT GTG GTG GGT-3' (SEQ ID NO:71) (an antisense primer corresponding to the sequence of positions 555 to 576 in SEQ derivative coding for positions 1 to 163 amino acid residues);

hTPO-13: 5'-AGT AAA CTG CTT CGT GAC TCC CAT (positions 124 to 173 in SEQ ID NO:4; prepared for use in the preparation of a derivative in which amino acid residues of GTC CTT CAC AGC AGA CTG AGC CAG TG-3" (SEQ ID NO:72) positions 1 to 12 are deleted);

64 to 87 and 124 to 148 in SEQ ID NO: 4, prepared for use in the IGG ACA GCG TTA GCC TTG CAG TTA G-3' (SEQ ID NO:73) (an antisense primer corresponding to the sequence of positions preparation of a derivative in which amino acid residues of hTPO-13R: 5'-CAT GGG AGT CAC GAA GCA GTT TAC positions 1 to 12 are deleted);

(positions 106 to 154 in SEQ ID NO:4, prepared for use in the preparation of a derivative in which amino acid residues of hTPO-7: 5'-TGT GAC CTC CGA GTC CTC AGT AAA CTG CTT CGT GAC TCC CAT GTC CTT C-3' (SEQ ID NO:74) positions 1 to 6 are deleted);

hTPO-7R: 5'-TTT ACT GAG GAC TCG GAG GTC ACA GGA CAG CGT TAG CCT TGC AGT TAG-3' (SEQ ID NO:75) (an antisense primer corresponding to the sequence of positions 64 to 87 and 106 to 129 in SEQ ID NO:4, prepared for use in the preparation of a derivative in which amino acid residues of positions 1 to 6 are deleted);

hTPO-8: 5'-GAC CTC CGA GTC CTC AGT AAA CTG positions 109 to 157 in SEQ ID NO:4, prepared for use in the CTT CGT GAC TCC CAT GTC CTT CAC A-3' (SEQ ID NO:76)

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performed in 100 µl volume making use of GeneAmpTM PCR System 9600 (manufactured by PERKIN-ELMER) and, after 5 minutes of denaturation at 95 c, repeating a total of 30 cycles, each cycle consisting of denaturation at 95°c for 1 minute, annealing at 60°c for 1 minute and synthesis at 72°c for 1 minute, followed by the final incubation at 72°c for 7 minutes. Each of the PCR products was subjected to 1.2% electrophoresis to isolate a main DNA fragment having an Gene DNA Purification Kit (manufactured by Bio-Rad) and subjected to extraction with the same volume of plasmid DNA of a deletion derivative (pHT13-231) coding for a gel (manufactured by FMC BioProducts) expected size which was subsequently purified using Prep-Adissolved in 15 µl of TE buffer. After digestion with restriction enzymes EcoRI and NotI, the resulting solution was phenol/chloroform and subsequent ethanol precipitation. After centrifugation, the resulting precipitate was dissolved in 15 ul of TE buffer and then sub-cloned into the expression vector pEF18S which has been treated in advance with the same restriction, enzymes. In this case, Competent-High E. coli DH5 manufactured by TOYOBO was used as the host strain. From the expected size were selected to prepare plasmid DNA samples basically in accordance with the procedure described in Molecular Cloning (Sambrook et al., Cold Spring Harbor to 12 have been deleted from the original amino acid residues of positions 1 to 231. When these 45 clones were subjected to these, the sequence of 3 clones were checked by 373A DNA resulting transformants, 45 clones containing the insert of Laboratory Press, 1989). In this way, it was able to obtain protein molecule in which amino acids residues of positions 1 Sequencer manufactured by Applied Biosystems making use of PCR using primers hTPO-5 and hTPO3, inserts having aq Dye DeoxyTM Terminater Cycle Sequening Kit (Applied approximately expected sizes were confirmed in 8 clones. agarose

Biosystems), thereby obtaining a clone pHT13-231 #3 having a TPO cDNA sequence as designed and a deletion as expected with no substitutions and the like over the entire nucleotide sednence.

(2) Preparation of a derivative (pHT7-163) in which amino acid residues of positions 1 to 6 are deleted

because, though the 231 position amino acid was designed as derivativess in the above step (1), it was found that TPO activity could be expressed even when the C-terminal amino In this case, the 163 position amino acid was used as the C-terminal for the preparation of deletion derivativess the C-terminal of TPO protein for the preparation of deletion the 163 position amino acid was used as the C-terminal also in plasmid DNA of the clone pHT1-163 obtained in Example 27 as rR in another combination) as primers. Using GeneAmpTM PCR Reagent Kit with AmpliTaqTM DNA Polymerase (manufactured manufactured by PERKIN-ELMER) and, after 5 minutes of (manufactured by FMC BioProducts) electrophoresis to isolate acids were further deleted. On the basis of the same reason, the subsequent step (3). PCR was carried out using 1.4 µg of a template and 5 µM each of the synthesized oligonucleotides (hTPO-7 and hTPO-S in one combination and hTPO-5 and hTPOby Takara Shuzo), the PCR reaction was performed in 100 μl volume making use of GeneAmpTM PCR System 9600 denaturation at 95°c, repeating a total of 30 cycles, each Each of the PCR products was subjected to 1.2% agarose gel a main DNA fragment having an expected size which was subsequently purified using Prep-A-Gene DNA Purification Kit cycle consisting of denaturation at 95°c for 1 minute, annealing at 65 c for 1 minute and synthesis at 72 c for 1 minute, followed by the final incubation at 72 c for 7 minutes. manufactured by Bio-Rad) and dissolved in 15 µl of TE buffer.

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Thereafter, second PCR was carried out using 1 µl portion each of the thus prepared solution as a template.

plasmid DNA of a deletion derivative (pHT7-163) coding for a The second PCR was carried out using 5 µM each of the synthesized primers (hTPO-5 and hTPO-S). Using GeneAmpTM PCR Reagent Kit with AmpliTaqTM DNA Polymerase (manufactured by Takara Shuzo), the PCR reaction was berformed in 100 μl volume making use of GeneAmpTM PCR System 9600 (manufactured by PERKIN-ELMER) and, after 5 minutes of denaturation at 95°c, repeating a total of 30 minute, annealing at 60 c for 1 minute and synthesis at 72 c or 1 minute, followed by the final incubation at 72°c for 7 minutes. Each of the PCR products was subjected to 1.2% agarose gel (manufactured by FMC BioProducts) electrophoresis to isolate a main DNA fragment having an Gene DNA Purification Kit (manufactured by Bio-Rad) and dissolved in 15 µl of TE buffer. After digestion with restriction enzymes EcoRI and Notl, the resulting solution was subjected to extraction with the same volume of phenol/chloroform and subsequent ethanol precipitation. After centrifugation, the resulting precipitate was dissolved in 15 ul of TE buffer and then sub-cloned into the expression vector DEF18S which has been treated in advance with the same restriction enzymes. In this case, Competent-High E. coli DH5 manufactured by TOYOBO was used as the host strain. From the resulting transformants, 30 clones containing the insert of expected size were selected to prepare plasmid DNA samples basically in accordance with the procedure described in Molecular Cloning (Sambrook et al., Cold Spring Harbor aboratory Press, 1989). In this way, it was able to obtain protein molecule in which amino acids residues of positions 1 to 6 have been deleted from the original amino acid residues of cycles, each cycle consisting of denaturation at 95°C for 1 expected size which was subsequently purified using Prep-A-

positions 1 to 163 in SEQ ID NO: 4. When these clones were #4 and #29, each having a TPO cDNA sequence as designed and subjected to PCR using primers hTPO-5 and hTPO-S, inserts naving approximately expected sizes were confirmed in all Of these, the sequences of 3 clones were checked by 373A DNA Sequencer manufactured by Applied Biosystems making use of Taq Dye DeoxyTM Terminater Cycle Sequening Kit Applied Biosystems), thereby obtaining 2 clones, pHT7-163 a deletion as expected with no substitutions and the like over he entire nucleotide sequence. clones.

amino acid residues of positions 1 to 7 are deleted Preparation of a derivative (pHT8-163) in which

acid residues of positions 1 to 7 was prepared basically in the same manner as the above case for the preparation of a 8R in another combination) as primers. Using GeneAmp TM PCR Reagent Kit with AmpliTaq TM DNA Polymerase (manufactured manufactured by PERKIN-ELMER) and, after 5 minutes of A derivative (pHT8-163) having deletion of amino derivative (pHT7-163) having deletion of amino acid residues a template and 5 µM each of the synthesized oligonucleotides (hTPO-8 and hTPO-S in one combination and hTPO-5 and hTPOby Takara Shuzo), the PCR reaction was performed in 100 µl volume making use of GeneAmpTM PCR System 9600 manufactured by FMC BioProducts) electrophoresis to isolate a main DNA fragment having an expected size which was of positions 1 to 6. PCR was carried out using 1.4 µg of denaturation at 95°c, repeating a total of 30 cycles, each annealing at 65°c for 1 minute and synthesis at 72°c for 1 ninute, followed by the final incubation at 72 c for 7 minutes. Each of the PCR products was subjected to 1.2% agarose gel ubsequently purified using Prep-A-Gene DNA Purification Kit cycle consisting of denaturation at 95°c for 1 minute, plasmid DNA of the clone pHT1-163 obtained in Example 27

Thereafter, second PCR was carried out using 1 µl portion of manufactured by Bio-Rad) and dissolved in 15 µl of TE buffer.

each of the thus prepared solutions as a template.

The second PCR was carried out using 5 µM each of

the synthesized primers (hTPO-5 and hTPO-S). Using GeneAmpTM PCR Reagent Kit with AmpiiTaqTM DNA Polymerase manufactured by Takara Shuzo), the PCR reaction was performed in 100 µl volume making use of GeneAmpTM PCR System 9600 (manufactured by PERKIN-ELMER) and, after 5 minutes of denaturation at 95 c, repeating a total of 30 minute, annealing at 60 c for 1 minute and synthesis at 72 c

ycles, each cycle consisting of denaturation at 95°c for 1

minutes. Each of the PCR products was subjected to 1.2%

or 1 minute, followed by the final incubation at 72°c for 7

electrophoresis to isolate a main DNA fragment having an

expected size which was subsequently purified using Prep-A-

Gene DNA Purification Kit (manufactured by Bio-Rad) and Jissolved, in 15 µl of TE buffer. After digestion with restriction enzymes EcoRI and NotI, the resulting solution was subjected to extraction with the same volume of phenol/chloroform and subsequent ethanol precipitation. After

agarose gel (manufactured by FMC BioProducts)

centrifugation, the resulting precipitate was dissolved in 15

ul of TE buffer and then sub-cloned into the expression vector

EF18S which has been treated in advance with the same

restriction enzymes. In this case, Competent-High E. coli DH5 nanufactured by TOYOBO was used as the host strain. From the expected size were selected to prepare plasmid DNA samples pasically in accordance with the procedure described in Molecular Cloning (Sambrook et al., Cold Spring Harbor aboratory Press, 1989). In this way, it was able to obtain plasmid DNA of a deletion derivative (pHT8-163) coding for a

esulting transformants, 30 clones containing the insert of

protein molecule in which amino acids residues of positions 1

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Expression of deletion derivative in COS cells and confirmation of TPO activity

Transfection of COS 1 cells with each of the thus obtained deletion clones was carried out according to the dextran method which includes chloroquine treatment, and the procedure of Example 11. That is, transfection was performed culture supernatants were recovered after 3 days of culture. The thus obtained culture supernatant was thoroughly dialyzed against the IMDM culture medium described in the foregoing with 10 µg of each of the plasmid DNA samples by the DEAEand evaluated by the M-07e assay system.

was detected in the culture supernatant of COS 1 cells ransfected with a clone coding for the deletion derivative consisting of 7 to 163 position amino acids. However, no TPO activity was detected in the culture supernatant of COS 1 cells ransfected with a clone coding for the deletion derivative As a result, weak, but significant, TPO activity consisting of 8 to 163 position amino acids or of 13 to 231 osition amino acids.

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An expression vector having all amino acid coding regions of human TPO cDNA presumed in SEQ ID NO:6 was constructed for the expression in mammalian cells. PCR was carried out in the following manner in order to prepare a DNA fragment which covers all human TPO cDNA coding regions.

Nucleotide sequences of the used primers are as follows: hTPO-I: 5'-TTG TGA CCT CCG AGT CCT CAG-3' (SEQ ID NO:78) (positions 105 to 125 in SEQ ID NO:6);

SA: 5'-CAG GTA TCC GGG GAT TTG GTC-3' (SEQ ID NO:79) (an antisense primer corresponding to the sequence of positions 745 to 765 in SEQ ID NO:6);

hTPO-P: 5'-TGC GTT TCC TGA TGC TTG TAG-3' (SEQ ID NO:80) (positions 503 to 523 in SEQ ID NO:6); and

adding a restriction enzyme Notl recognition sequence and a corresponding to the sequence of positions 1066 to 1086 in htpo-ko: 5-GAG AGA GCG GCC GCT TAC CCT TCC IGA GAC. AGA TT-3' (SEQ ID NO:81) (a sequence prepared by GAGAGA sequence (SEQ ID NO:82) to an antisense sequence SEQ ID NO:6).

cycle consisting of incubations at 96 c for 1 minute, at 62 c Using 0.5 µM each of the primers hTPO-I and SA and 1 unit of Vent RTM DNA polymerase (manufactured by New England or 1 minute and at 72 c for 1 minute, followed by the final as follows: 10 mM KCl, 10 mM (NH4) $_2$ SO $_4$, 20 mM Tris-HCl (pH A first PCR was carried out using 300 ng of the BioLabs), the PCR (repetition of a total of 30 cycles, each Composition of the reaction solution in final concentration is clone pEF18S-HL34 obtained in Example 16 as a template. ncubation at 72.c for 7 minutes) was carried out. 3.8), 2 mM MgSO₄, 0.1% Triton X-100 and 200 μ M dNTP mix.

manufactured by Clontech) was heated at 70 c for 10 SuperScript TM II RNase H (manufactured by LIFE TECHNOLOGIES), followed by 1 hour of incubation at 37 c to of a total of 30 cycles, each cycle consisting of incubations at 96.c for 1 minute, at 58.c for 1 minute and at 72.c for 1 minute) was carried out using 1/20 volume of the reaction One microgram portion of commercially available ooly(A)+ RNA preparation derived from human normal liver minutes, rapidly cooled on ice and then mixed with 10 mM of DTT, 500 µM of dNTP mix, 25 ng of random primer manufactured by Takara Shuzo), 10 units of RNase Inhibitor manufactured, by Boehringer-Mannheim) and 200 units of perform synthesis of cDNA. Thereafter, second PCR (repetition solution of synthesized cDNA as a template and 2.5 µM each of the primers hTPO-P and hTPO-KO and 2.5 units of AmpliTaqTM DNA polymerase (manufactured by Takara Shuzo).

to isolate, respective main DNA fragments of expected size Purification Kit (manufactured by Bio-Rad). Thereafter, third oCR was carried out using 1/20 volume of each of the thus ninute, and then incubated at 72°c for 7 minutes. The Each of the resulting solutions of the first and second PCR was subjected to 1% agarose gel electrophoresis which was subsequently purified using Prep-A-Gene DNA prepared solutions as a template. This reaction (heating at 36.c for 2 minutes, followed by the repetition of a total of 3 cycles, each cycle consisting of incubations at 96 c for 2 minute and at 72.c for 2 minute, and subsequent incubation at 72.c for 7 minutes) was carried out using 1 unit of Vent $\mathsf{R}^\mathsf{T}\mathsf{M}$ DNA polymerase (manufactured by New England BioLabs). The esulting reaction solution was mixed with 1 µM each of hTPOl and hTPO-KO, heated at 96 c for 2 minutes, subjected to 25 cycles of reaction, each cycle consisting of incubations at 36.c for 1 minute, at 62.c for 1 minute and at 72.c for 1 esulting reaction solution was extracted with the same

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manufactured by Boehringer-Mannheim) to recover DNA. The the same volume of chloroform, and the extract was subjected to ethanol precipitation (2.5 volumes of ethanol in the presence of 0.3 M sodium acetate and 0.5 µl of glycogen electrophoresis, and a main band having an expected size thus volume of water saturated phenol-chloroform and then with hus recovered DNA was digested with restriction enzymes BamHI and NotI and subjected to 1% agarose gel isolated was purified using Prep-A-Gene DNA Purification Kit manufactured by Bio-Rad), ligated with pBluescript II SK+ vector (manufactured by Stratagene) which had been digested then transformed into Competent-High E. coli DH5 (manufactured by TOYOBO). From the resulting colonies, 4 clones were selected to prepare plasmid DNA samples. The sequence of purified plasmid DNA samples were checked by 373A DNA Sequencer manufactured by Applied Biosystems naking use of Taq Dye DeoxyTM Terminater Cycle Sequening Kit (Applied Biosystems), thereby obtaining a clone, pBLTP, having a TPO cDNA sequence as designed with no substitution in n advance with the restriction enzymes BamHI and Notl and nucleotide sequence within the region between BamHI and Notl.

The clone pBLTP was digested with restriction enzymes EcoRI and BamHI and subjected to 1% agarose gel electrophoresis, and a band of high molecular weight thus isolated was purified using Prep-A-Gene DNA Purification Kit was treated with the restriction enzymes to purify a DNA ragment of about 450 bp. Each of the thus obtained DNA samples was subjected to ligation and transformed into Plasmid DNA samples were prepared from the resulting manufactured by Bio-Rad). In the same manner, pEF18S-HL34 colonies to obtain a clone, pBLTEN, containing human TPO cDNA The thus obtained pBLTEN was digested with estriction enzymes EcoRI and NotI and subjected to 1% Competent-High E. coli DH5 (manufactured by TOYOBO),

garose gel electrophoresis, and a DNA fragment of about 1,200 bp thus isolated was purified using Prep-A-Gene DNA Purification Kit (manufactured by Bio-Rad), ligated with the expression vector pEF18S which has been digested in advance with the same restriction enzymes and then transformed into Plasmid DNA samples were prepared from the resulting colonies to obtain the clone of interest, pHTP1, which contains entire human TPO cDNA coding region. Plasmid DNA was prepared from this clone in a large quantity and used in the ollowing experiments. Preparation of plasmid DNA was described in Molecular Cloning. (Sambrook et al., Cold Spring Competent-High E. coli DH5 (manufactured by TOYOBO). carried out basically in accordance with the procedure Harbor Laboratory Press, 1989).

Example 31>

Construction of mammalian expression plasmid, pDEF202hTPO-P1, for the expression of human TPO in Chinese hamster ovary (CHO) cells.

One microgram of a plasmid pMG1 containing mouse estriction enzymes, EcoRI and BamHI, and the subjected to fragment was dissolved in 25 ml of a reaction solution composed of 50 mM Tris-HCl (pH 7.5), 7 mM MgCl $_{
m 2}$, 1 mM 2mercaptoethanol and 0.2 mM dNTP mixture, mixed with 2 units dihydrofolate reductase (DHFR) minigene was digested with agarose gel electrophoresis to isolate a fragment (about 2.5 containing mouse DHFR minigene. The isolated DNA of Klenow fragment and then incubated at room temperature for 30 miniutes to form blunt ends at both termini of the DNA ragment. After phenol/chloroform treatment and ethanol precipitation, the resulting fragment was dissolved in 10 ml of a TE solution (10 mM Tris-HCI (pH 8.0)/1 mM EDTA). The nammalian expression vector pEF18S was digested with a estriction enzyme Smal, dephosphorylated with alkaline

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phosphatase (manufactured by Takara Shuzo) and then ligated with a DNA fragment containing mouse DHFR minigene by using T4 DNA ligase (manufactured by Takara Shuzo) to obtain an expression vector pDEF202.

Next, the constructed expression vector pDEF202 then subjected to agarose gele electrophoresis to isolate a arger DNA fragment. Human TPO cDNA which has been obtained by digesting the plasmid pHTP1 containing human TPO cDNA (clone p1) with restriction enzymes, EcoRI and Spel, was early was digested with restriction enzymes, EcoRI and Spel, and DEF202-hTPO-P1 for expression of human TPO cDNA. This constructed plasmid contains SV40 replication origin, human igated with this linearized pDEF202 vector by using T4 DNA igase (manufactured by Takara Shuzo) to obtain a plasmid polyadenylation signal for transcription of human TPO cDNA, mouse DHFR minigene, and pUC18-replication origin and β and SV40 factor 1-a-promoter, actamase gene (Amp^r) elongation

<Example 32>

Expression of human TPO in CHO cells

CHO cells (a DHFR- strain; Urlaub and Chasin, Proc. Natl. Acad. Sci. USA, vol.77, p4216, 1980) were maintained in α-minimum essential medium (α-MEM(-), supplemented with hypoxanthine and thymidine) containing 10% fetal calf serum (FCS) by using a 6 cm diameter tissue culture plate (by Falcon). The transformation of CHO cells with a pDEF202-hTPO-P1 plasmid was carried out by a calcium phosphate method (CellPhect, manufactured by Pharmacia) as described below. Ten micrograms of the plasmid pDEF202-hTPO-P1 prepared in Example 31 was mixed with 120 ml of buffer A and 120 ml of H2O, and the resulting mixture was incubatedat room temperature for 10 minutes. This solution was further mixed with 120 ml of buffer B and allowed to stand at room

solution was then added into the culture and incubated for 6 emperature for additional 30 minutes. This DNA mixture hours in a CO2 incubator. After 6 hours, the culture was MEM(-) containing 10% dimethyl sulfoxide for 2 minutes, and supplemented with hypoxanthine and thymidine) containing 10 For selection of transformed cells with DHFR-positive phenotype, the cells were treated with a trypsin/EDTA in 6 cm tissue culture plate were split into five 10 cm tissue washed twice with serum-free α -MEM(-), treated with α then incubated for 2 days in non-selection medium (a-MEM(-) % dialyzed FCS. After 2 days culture, the cells were selected in a selection medium (α-MEM(-)) containing 10% dialyzed FCS. solution, and resuspended with a selection medium. The cells culture plates or twelve 24-well tissue culture plates, and then cultured in the selection medium. The culture medium was exchanged at the intervals of two days. The human TPO activity in the cultured medium of CHO cells with DHFRpositive phenotype was measured by CFU-MK assay, M-O7e assay or. Ba/F3 assay. The cells secreting human TPO into culture medium were further selected in the selection medium supplemented with 25 nM methotrexate in order to isolate the sell clone producing a higher amount of human TPO.

In this instance, transfection of CHO cells may also be effected by carrying out co-transfection of CHO cells with pHTP1 and pMG1 plasmids.

CHO strain (CHO-DUKXB11) transfected with the plasmid pDEF202-hTPO-P1 has been deposited by the present applicant on January 31, 1995, under the accession No. FERM BP-4988, at the National Institute of Bioscience and Human Technology, Agency of Industrial Science and Technology, Ministry of International Trade and Industry, Japan.

cExample 33>

Construction of a recombinant vector,

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BMCGSneo-hTPO-P1, for use in X 63.6.5.3. cells

Not!, and the subjected to agarose gel electrophoresis to solate the vector DNA portion. This DNA fragment and human rPO cDNA (P1 clone) which had been obtained by digesting the plasmid pBLTEN containing human TPO cDNA with the pBMCGSneo was digested with restriction enzymes, Xhol and estriction enzymes Xhol and Notl were ligated using T4 DNA igase to obtain the expression vector of interest, pBMCGSneohTPO-P1. This expression plasmid contains cytomegalovirus early promoter, parts of rabbit \$ globin gene-derived intron and polyadenylated region for transcription of human TPO cDNA, human ß globin gene, 69% of bovine papilloma virus 1 phosphotransferase I gene (neomycin resistant gene) and pBR One microgram of a mammalian expression vector gene, thymidine kinase promoter and polyadenyl region, 322 replication origin and β-lactamase gene (Amp^r), and human TPO cDNA was ligated at a downstream site of the cytomegalovirus promoter.

Example 34>

Expression in X 63,6.5.3 cells

minimal essential (DME) medium containing 10% fetal bovine serum. The transfection of X63.6.5.3 cells with BMCGSneo hTPO X 63.6.5.3.cells were cultured in Dulbecco's plasmid was carried out by electroporation method as described below.

prepared in Example 33 were added to 750 µl of the DME electroporation cuvette having a gap of 4 mm and allowed to electroporation apparatus (BTX 600, manufactured by BTX) to carry out gene transfer under conditions of 380 V, 25 mF and medium containing 107 cells, and the mixture was put into an stand for 10 minutes at 4.c. The cuvette was then set into an 24 angstroms. After the gene transfer, the cuvette was again 20 micrograms of the plasmid BMCGSneo-hTPO-P1

in a CO2 incubator. After the 2 days culture, the culture was mesured by CFU-MK, M-07e or Ba/F3 assay. The ransformants secreting human TPO into culture medium were allowed to stand for 15 minutes at 4.c, and the cells were the transfected cells were suspended in 50 ml of DEM containing 10% fetal bovine serum, dispensed into wells of ive 96 well tissue culture plate and then cultured for 2 days medium was exchanged with DEM containing 10% fetal bovine serum and 1 mg/ml of G418 (manufactured by GIBSCO), and the nedium exchange was carried out every 3 days thereafter. The human TPO activity in the culture medium of transformants cloned twice by a limiting dilution to establish a human TPO then washed once with DEM containing 10% fetal bovine serum. producing cell lines.

<Example 35>

Large scale expression of human TPO in COS 1 cells

FCS using a collagen-coated 175 cm² culture flask at 37°C in a according to the DEAE-dextran method which includes chloroquine treatment as described in Example 11. COS 1 cells 5% carbon dioxide incubator until the cells became about 100% confluent. In order to effect collagen coating of the culture manufactured by Iwaki) whose concentration has been adjusted flask was washed once with 20 to 50 ml of PBS. Transfection was effected by mixing 20 ml of IMDM solution containing 250 chloroquine (manufactured by Sigma) and 10% (v/v) of Nu-Transfection of COS 1 cells was carried out (ATCC CRL1650) were cultured in IMDM containing 10% (v/v) flask, 25 ml of a collagen solution (Cellmatrix type I-C, to 0.3 mg/ml with 1 mM HCl was added to each 175 cm² culture flask, the collagen solution was recovered after 1 hour of standing at room temperature and then the thus treated μg/ml of DEAE-dextran (manufactured by Pharmacia), 60 μM of Serum (manufactured by Collaborative) with the plasmid

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vith 50 ml of a serum-free medium and then cultured for 5 vashed once with IMDM just before the transfection, and then sulturing the cells for 3 hours at 37 c in a 5% carbon dioxide suction, and the cells were washed once with IMDM, mixed days at 37°c in the 5% carbon dioxide incubator to recover the culture supernatant. In one operation, 100 to 260 culture lasks of 175 cm² were used and 5 to 13 liters of the culture supernatant was recovered. The serum-free medium was prepared by supplementing IMDM with 5 µg/ml of insulin manufactured by Sigma), 5 µg/ml of transferrin manufactured by Sigma), 10 µM of monoethanolamine manufactured by Wako Pure Chemical Industries), 25 nM of sodium selenite (manufactured by Sigma) and 200 µg/ml of bHTP1 (40 μg) which has been dissolved in 500 μl of HBS, adding the resulting mixture to the COS 1 cells described bove contained in one 175 cm² culture flask, which have been ncubator. Thereafter, the culture supernatant was removed by BSA (fatty acid-free high purity bovine albumin, manufactured by Nichirei).

Example 36>

Purification of human full-length TPO (derived from expression plasmid pHTP1. P1 clone) expressed in COS 1 cells and its biological activity

activity in the following preparation steps. TPO activity was vas carried out to obtain a high purity product. The M-07e assay system was mainly used for the measurement of TPO and purification of human full-length TPO (derived from the expression plasmid pHTP1) expressed in COS 1 cells. A partially purified product was firstly prepared in order to examine platelet increasing activity and then the purification similarly found in the rat CFU-MK assay system. In carrying The following déscribes examples of the activity

out these assays, human serum albumin (HSA) was added to each sample to a final concentration of 0.02 to 0.05%. Firstly, COS 1 cells transfected with the plasmid HTP1 according to the method of Ohashi and Sudo (Hideya Chashi and Tadashi Sudo, Biosci. Biotech. Biochem., 58 (4), 758 759, 1994) were cultured for 5 days at 37 c in a 5% CO2 using a serum-free IMDM culture medium containing, 0.2 g of 3SA, 5 mg of bovine insulin, 5 mg of human transferrin, 0.02 nM of monoethanolamine and 25 nM of sodium selenite, in 1000ml, thereby obtaining about 7 liters of serum-free culture supernatant. To the serum-free culture supernatant was added Millipore), 723 ml of the thus concentrate (protein concentration, 3.38 mg/ml; total protein, 2,445 mg; relative proteolytic enzyme inhibitors p-APMSF and Pefabloc SC (4-(2manufactured by Merk, catalog No. 24839) to a final concentration of about 1 mM, followed by filtration using a 0.2 um filter to recover the supernatant. This was concentrated by a factor of about 10 using an ultrafiltration unit (Omega with 1.6 moles per 1,000 ml of ammonium sulfate (288 g in aminoethyl)-benzenesulfonyl fluoride hydrochloride, manufactured by Filtron; or PLGC Pellicon Cassette, nominal molecular weight cutoff of 10,000, manufactured by activity, 43,000; and total activity, 105,100,000), was mixed total) to obtain 804 ml of the solution containing 1.5 M in final concentration of ammonium sulfate, and then the thus obtained nas been equilibrated in advance with 20 mM sodium citrate buffer (pH 5.2) containing 1.5 M ammonium sulfate. After the solution was applied at a flow rate of 10 ml/min to Macro-Prep Methyl HIC column (5 cm in diameter and 9 cm in bed height, manufactured by Bio-Rad, catalog No. 156-0080) which sample loading, elution was carried out with 20 mM sodium citrate buffer (pH 5.2) containing 1.25 M ammonium sulfate to obtain passed-through fraction F1 (2,384 ml; protein Ultrasette, nominal molecular weight cutoff of 8,000,

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concentration, 0.864 mg/ml; total protein, 2,061 mg; and relative activity, 6,000).

Next, the eluting solution was changed to 20 mM Na citrate, pH 5.8, to collect fraction F2 (1,092 ml; protein concentration, 0.776 mg/ml; total protein, 847 mg; and relative activity, 150,000).

The thus obtained Macro-Prep Methyl HIC column raction F2 (1,081 ml) was applied at a flow rate of 10 ml/min to SP Sepharose Fast Flow column (3 cm in diameter and 10 cm in bed height, manufactured by Pharmacia Biotech, catalog No. 17-0729-01) which has been equilibrated in advance with 20 nM sodium citrate buffer (pH 5.8). After the sample loading, elution was carried out with 20 mM sodium citrate buffer (pH protein concentration, 0.270 mg/ml; total protein, 610 mg; and relative activity, 30,000). Next, the eluting solution was 100 mM NaCl to collect fraction F2 (856 ml; protein concentration, 0.189 mg/ml; total protein, 162 mg; and elative activity, 300,000). Next, the eluting solution was 5.6) containing 110 mM NaCl to obtain fraction F1 (2,262 ml; changed to 20 mM sodium citrate buffer (pH 5.4) containing changed to 20 mM sodium citrate buffer (pH 5.2) containing ,000 mM NaCl to collect eluted fraction F3 (370 ml; protein concentration, 0.034 mg/ml; total protein, 12.6 mg; and elative activity, 150,000).

The main TPO active fraction F2 (845 ml) of the SP Sepharose Fast Flow column step was mixed with about 10% in final concentration of 1-propanol and applied at a flow rate of 3 ml/min to LA-WGA column (2 cm in diameter and 14 cm in bed height, manufactured by Honen, catalog No. WG-007) which has been equilibrated in advance with 20 mM sodium citrate buffer (pH 5.4) containing 400 mM NaCl. After the sample loading, elution was carried out using a mixture of 20 mM sodium citrate buffer (pH 5.4) containing 400 mM NaCl. After the sample propanol (9:1) to obtain fraction F1 (64.4 ml; protein

concentration, 0.0178 mg/ml; total protein, 1.15 mg; and relative activity, 17,220). Next, the eluting solution was changed to 20 mM sodium citrate buffer (pH 6.1) containing 0.4 M GlcNAc and 10% 1-propanol to collect eluted fraction F2 (45 ml; protein concentration, 0.0104 mg/ml; total protein, 0.470 mg; and relative activity, 675,000).

The main TPO active fraction F2 (340 ml) of the LA-WGA column operation was mixed with about 0.005% in final concentration of TFA and subjected to YMC-Pack CN-AP (6 mm in diameter and 250 mm in bed height, manufactured by YMC, (catalog No. AP-513) column chromatography. That is, using 0.1% TFA as a developing solvent A and 1-propanol containing 0.05% TFA as a developing solvent B, the sample was applied at a flow rate of 0.6 ml/min to the YMC-Pack CN-AP column which has been equilibrated in advance with 15% B. After the sample loading, the propanol concentration was increased from 15% B to 25% B, and the elution was carried out with a linear gradient of from 25% B to 50% B for 65 minutes while collecting the eluates in 1.5 ml (2.5 min) portions in polypropylene tubes.

A 0.5 µl portion (1/3,000 fraction) of the eluate in each tube was mixed with HSA and concentrated by ultrafiltration to obtain 0.25 ml of IMDM assay culture solution containing 0.05% HSA for use in the identification of TPO active fractions by the assay. As the result, strong TPO activity (relative activity of about 630,000 to 4,800,000) was found in tube Nos. 24 to 30 (within the range of 35.0 and 42.0% as propanol concentration) which were pooled as TPO active fraction FA (13.5 ml).

The fraction FA obtained by the YMC-Pack CN-AP column step was subjected to Capcell Pak Cl 300A (4.6 mm in diameter and 150 mm in bed height, manufactured by Shiseido, catalog No. Cl TYPE:SG300A) column chromatography using 0.1% TFA as a developing solvent A and 1-propanol containing 0.05%

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for 5 minutes and then with a linear gradient of from 20% B to A portion (8.9 ml) of the a flow rate of 0.4 ml/min to the Capcell Pak Cl 300A column which has been equilibrated in advance with 20% B. After the sample loading, the elution was carried out firstly with 20% B 40% B for 50 minutes while collecting the eluates in 1 ml (2.5 was mixed with 0.3 ml of glycerol, concentrated by centrifugation evaporation and then mixed with about 2.5 ml of 10% B, and the resulting solution was subsequently applied at raction FA obtained by the YMC-Pack CN-AP column operation min) portions in polypropylene tubes. IFA as a developing solvent B.

A 1 µl portion (1/1,000 fraction) of the eluate in ultrafiltration to obtain 0.25 ml of IMDM assay culture TPO active fractions by the assay. As the result, strong TPO each tube was mixed with HSA and concentrated by solution containing 0.05% HSA for use in the identification of was found in tube Nos. 20 to 23 (within the range of 28.5 and activity (relative activity of about 3,000,000 to 22,500,000) 32.5% as propanol concentration) which were pooled as a high activity fraction.

<Example 37>

Molecular weight measurement of human TPO expressed in COS 1 cells

the human complete length TPO (derived from the expression It was assumed that molecular weight of the human complete length TPO (derived from the expression plasmid pHTP1, F1 clone) expressed in COS 1 cells would be larger than the motecular weight presumable from the size of the peptide as a first step, a partially purified TPO fraction was prepared in the following manner from a culture supernatant containing chain because of the addition of sugar chain. In consequence, plasmid pHTP1, F1 clone) expressed in COS 1 cells. That is, using a developing solvent A (0.1% trifluoroacetic acid (TFA))

a 0.3 ml portion of the culture supernatant was applied to rate of 0.4 ml/min and then fractionation was effected by a inear density gradient of from 25% B to 50% B spending 50 ninutes. TPO activity was found in eluates within the range of 34.5% and 43.5% as propanol concentration, and these eluates were evaporated to dryness by centrifugation evaporation to YMC-Pack PROTEIN-RP (0.46 cm in diameter and 15 cm in bed column which has been equilibrated in advance with 25% B, 25% B was passed through the column for 5 minutes at a flow neight, manufactured by YMC, catalog No. A-PRRP-33-46-25) and a developing solvent B (1-propanol containing 0.05% TFA), obtain a partially purified sample.

Next, TPO activity was examined by the M-07e assay system after extracting protein from a gel of SDS-PAGE electrophoresis carried out under a non-reducing condition as pased on reduction-treated DPCIII markers by means of a reated DPCIII markers after its N-glycosidic linkage type sugar chain digestion with N-glycanase (manufactured by Genzyme, catalog No. 1472-00) was measured to be 36,000 to 10,000 which was larger than the molecular weight of about 35,000 presumable from the size of the peptide chain, thus described in Example 1, or molecular weight was measured As a result, TPO activity was found within a broad apparent notecular weight range of approximately from 69,000 to 34,000, thus confirming variation in the molecular weight. Also, apparent molecular weight of TPO based on reductionstrongly indicating the presence of an O-glycosidic linkage western analysis which will be described later in Example 45. ype sugar chain too.

In the same manner, apparent molecular weight of the human complete length TPO (derived from the expression plasmid pHTP1, P1 clone) expressed in COS 1 cells was found - 176

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Example 38>

Biological characteristics of human IPO

Effects of human TPO on human and rat nematopoietic cells were examined, mainly using culture supernatant from COS 1 cells transfected with pHTF1.

In the colony assay using CD34+DR+ cell fraction prepared from human cord blood, human TPO stimulated the For example, 11.5 megakaryocyte colonies were formed by 5,000 CD34+DR+ cells in the presence of 10% (v/v) of culture supernatant from COS1 cells transfected with pHT1-231. To examine the effect of human TPO on human megakaryocyte progenitor cells in peripheral blood, CD34+ cells were prepared rom human peripheral blood as follows. A leukocyte fraction obtained from human peripheral blood by a Ficoll-Paque density gradient was depleted of cells adherent to plastic dishes. The nonadherent cells, subsequently, were depleted of SBA (soybean agglutinin) positive cells by a panning using AIS Microselecter SBA(Asahi Medical). The resultant cells were CD 34+ cells, were collected when CD3+ cells were cultured ior 10 days in the presence of culture supernatant from ransfected COS 1 cells in a liquid medium, selective proliferation of large size Gpllb/Illa+ cells was observed, and ncrease in the ploidy was found in these GpIIb/IIIa+ cells. these results strongly suggested that human TPO exerts its selective action on progenitor cells of human megakaryocyte ormation of significant number of megakaryocyte colonies. ncubated on AIS Microselecter CD 34, and the adsorbed cells, ineage and stimulates their proliferation and differentiation.

in the colony assay using GpIIb/IIIa+ cells highly enriched for CFU-MK from rat bone marrow cells and plastic nonadherent cells obtained at the preceding step of GpIIb/IIIa+ cells, human TPO induced the formation of a significant number of megakaryocyte colonies. For example, 34.5 megakaryocyte colonies were formed by 1,000 GpIIb/IIIa+

cells and 28.5 megakaryocyte colonies were formed by 20,000 plastic nonadherent cells in the presence of 20% (v/v) of culture supernatant from transfected COS 1 cells. In addition, although the plastic nonadherent cell fraction contains various hematopoietic progenitor cells other than CFU-MK, only megakaryocyte colonies were formed in the presence of human TPO, strongly suggesting that human TPO exerts its selective action on progenitor cells of the megakaryocyte lineage. When the GpIIb/IIIa⁺ cells obtained from rat bone marrow was cultured for 3 to 5 days in the presence of 20% (v/v) of culture supernatant from transfected COS 1 cells in a liquid culture, increase in the ploidy was observed clearly on 5 days of the

<Example 39>

Construction of vector for use in the expression of a fusion protein (to be referred to as "GST-TPO(1-174)" hereinafter) of glutathione-S-transferase (GST) and human TPO (amino acid residues 1, to 174) in E. coli

In order to facilitate expression of human TPO in *E.* coli, an artificial gene which encodes human TPO and contains an *E. coli* preference codon was prepared. The nucleotide sequence of this DNA contains an amino-terminal methionine codon (ATG) at the -1 position for use in the translation intitiation in *E. coli*.

Synthetic oligonucleotides 1 to 12:

1:5-CTAGAAAAACCAAGGAGGTAATAAATAATGAGOCC

GGCTCCGCCAGCTTGTGACCTTCGTGTTCTGTCT-3 (SEQ ID NO.83); 2.5.CAGTTTAGACAGAACGTCACAAGCTGGCCG

AGOGGGCTCATTATTATTACTCCTTGGTTTTTT-3' (SEO ID NO.84); S: 5-AAACTGCTTCGGAGCTCCACTCTCG TCTGTCCCAGTGCCCGGAAGTTCACCCGCTGCCG-3 (SEQ.ID.NOAS); \$ SCGGGGGTCGACTTCCGGGCCTGGG ACAGACGAGAGTGCAGCACGTGAGAGTCGCGAAG-3 (SEQ ID NOXB);

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6: 5-ACCOGETTICTECTTCCGCTGTCGACTTCTCCCT

GGGTGAATGGAAAACCCAGATGGAAGAGACCAAA-3 (SEQID NOB7); S. S-CTGAGCTTTGGTCTTCCATCTGGGTTTTCCATT

CACCCAGGGAGAAGTCGACAGCCGGAAGCAGAAC-3 (SECI ID NO88); 2: S-COTCAGGACATOCTGGGTGCAGTAACTCTGCTTCT GGAAGGCGTTATGGCTGCACGTGGCCAGCTTGGC3 (SEQ ID NO.89); B. S-GSTCGGGCCAAGCTGGCCACGTGCAGCCATAACGC

CTTCCAGAAGCAGAGTTACTGCACCCAGGATGTC-3 (SEQ ID NO:30); R SCOGAOCTECCTETTECCTECTTECCCAGCTETC rescondent to the respondence of IR SCAGAGACTTECAGAGOGOCGAGCAGAOGAAOCTTEGC CAGACAGCTGGCCAAGCAGGACAGGCA3 (SEQ ID NO.32); 11:5-TCTCTGCTTGGCACCAGCTGCCGCCACAGGGCCCG TACCACTECTCACAAGGATCCGAACGCTATCTTCC:3 (SEQ ID NO33); and 12:5-AAGACAGGAAGATAGCGTTCGGATCCTTGTGAGCA STEGTACGGCCCTGTGGCGGCCACCTGGGTGCCAAG-3 (SEQIDNOS4)

acetate, 10 mM Mg-acetate and 50 mM K-acetate. In order to nto combinations of 1 and 2, 3 and 4, 5 and 6, 7 and 8, 9 and 10 and 11 and 12, and each combination was subjected to were prepared and each of the synthetic oligonucleotides 2 to Pharmacia) in a solution composed of 1 mM ATP, 10 mM Trismake these synthetic oligonucleotides into 6 double-stranded DNA fragments, these synthetic oligonucleotides were made 10 mM MgCl₂ and 50 mM NaCl. Next, 3 double-stranded DNA 11 was phosphorylated with T4 kinase (manufactured by ragments of 1 and 2, 3 and 4, and 5 and 6 and the other 3 reated with T4 ligase in the same manner. The DNA fragment obtained by the ligation reaction was digested with BamHI annealing in a solution composed of 10 mM Tris/HCI (pH 7.5), double-stranded DNA fragments of 7 and 8, 9 and 10 and 11 and 12 were respectively treated with T4 ligase (manufactured by Life Technology), and these 2 reaction solutions were again manufactured by Boehringer-Mannheim) and subjected to 2%

was used as the host). From the clones thus obtained, a clone has been digested in advance with Xbal and BamHI (E. coli DH5lphahaving the artificial gene which encodes human TPO and contains the following E. coli preference codons was selected agarose gel electrophoresis to recover a fragment of about 390 to 400 bp in size which was subsequently purified using Prep-A-Gene DNA Purification Kit and sub-cloned into pUC18 which DNA sequence of the coding chain is shown in the Sequence by nucleotide sequence analysis and named pUC18(XB)(1-123). .isting (SEQ ID NO:9).

CTAGAAAAAA CCAAGGAGGT AATAAATAAT GAGCCCGGCT CCGCCAGCTT GTGACCTTCG ITITII GGITCCICCA ITAIITAITA CICGGCCGA GGCGGICGAA CACIGGAAGC 20 40 8 2 10 IE

TGITCIGICI AAACIGCIIC GOSACICICA CGIGCIGCAC ICICGICIGI CCCAGIGCCC ACAAGACAGA TITGACGAAG CGCTGAGAGT GCACGACGTG AGAGCAGACA GGGTCACGGG 8

110

100

8

180 GGAAGTICAC CCGCIGCCGA CCCCGGIICI GCTICCGGCI GICGACTICI CCCIGGGIGA CCITCAAGIG GGCGACGGCC GGGCCCAAGA CGAAGGCCGA CAGCIGAAGA GGGACCCACI 170 2 160 150 140 130

240 ATGGAAAACC CAGATGGAAG AGACCAAAGC TCAGGACATC CTGGGTGCAG TAACTCTGCT PACCITITIGG GICIACCIIC ICIGGITICG AGICCIGIAG GACCCACGIC AITGAGACGA 2 230 220 210 200 190

8 8 ICTGGAAGGC GTTATGGCTG CACGTGGCCA GCTTGGCCCG ACCTGCCTGT CTTCCCTGCT agacctrocs caatacceac stscaccest cgaaccesec tsgacgsala gaaggsacsa 290 280 270 260 250

resochecte reresoches ircsreter seresseser erscherer recifeche ACCGGTCGAC AGACCGGTCC AAGCAGACGA CGAGCCGCGA GACGTCAGAG ACGAACCGTG 350 340 330 320

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GGTCGACGGC GGTGTCCGG CATGGTGACG AGTGTTCCTA GGCTTGCGAT AGAAGGACAG AA 420 CCAGCTGCCG CCACAGGCC GTACCACTGC TCACAAGGAT CCGAACGCTA TCTTCC 410 400 330 380 4

95 c for 1 minute, at 58 c for 1 minute and at 72 c for 1 primer, and PCR was carried out using 0.1 volume of the cDNA minute, and final incubation at 72°c for 7 minutes) was Single-stranded cDNA was synthesized from 1 µg synthesis reaction solution as a template. The PCR reaction incubation at 96 c for 2 minutes, repetition of a total of 30 cycles of reaction, each cycle consisting of incubations at carried out with a volume of 100 µl using hTPO-C and hTPO-Z(EcoRI) as primers. The human TPO cDNA fragment obtained in this way was digested with BamHI and EcoRI and subjected to Gene DNA Purification Kit and sub-cloned into pUC18 which has been digested in advance with EcoRI and BamHI (E. coli DH5a was used as the host). A clone encoding correct human TPO nucleotide sequence was selected by nucleotide sequence analysis and named pUC18(BE)(124-332). Oligonucleotide sequences of the primers used in the PCR reaction are as of human normal liver-derived poly (A)+ RNA using oligo dT 2% agarose gel electrophoresis to recover a fragment of about 600 bp in size which was subsequently purified using Prep-A-

(SEQ ID NO:95) (positions 329 to 349 of pHTF1 clone); and hTPO-C: 5'-GGA GGA GAC CAA GGC ACA GGA-3'

recognition sequence to an antisense primer corresponding to hTPO-Z(EcoRI): 5-CCG GAA TTC TTA CCC TTC CTG AGA CAG ATT-3' (SEQ ID NO:96) (prepared by adding an EcoRI positions 1,143 to 1,163 of pHTF1 clone).

The clone pUC18(BE)(124-332) was digested with BamHI and EcoRI and subjected to 2% agarose gel

cloned into pUC18(XB)(1-123) which has been digested in advance with EcoRI and BamHI (E. coli DH5 α was used as the electrophoresis to recover a human TPO cDNA C-terminal side ragment of about 600 bp in size which was subsequently A clone obtained in this manner was named ourified using Prep-A-Gene DNA Purification Kit and sub-OC18(XE)(1-332). nost).

erminal deletion constructs of human TPO cDNA were prepared and expressed to measure in vitro human TPO Since the presence of human TPO activity in the residues had been confirmed by an example in which various Cactivity, an expression vector containing this peptide fragment numan TPO peptide fragment of 1 to 163 position amino acid was constructed. In this case, expression of the DNA sequence which encodes GST-TPO(1-174) was carried out.

Saci, two terminal codons were introduced using 2 synthetic oligonucleotides making use of the recognition site of this estriction enzyme. That is, two synthetic oligonucleotides erminal fragment of about 480 bp has been removed by its digestion with Sacl and EcoRI, thereby obtaining a clone Sequences of the synthetic oligonucleotides used herein are as Since a nucleotide sequence corresponding to positions 681 to 686 (amino acid residues of positions 173 and 174) of the pHTF1 clone is recognized by a restriction enzyme ris/HCl (pH 7.5), 10 mM MgCl₂ and 50 mM NaCl, and, making use of DNA Ligation Kit (manufactured by Takara Shuzo), the thus obtained double-stranded DNA fragment was introduced SSE1 and SSE2 were annealed in a solution composed of 10 mM nto pUC18(XE)(1-332) from which the human TPO cDNA C-O(18(XS)(1-174) (E. coli DH5 α was used as the host).

CTAATGAG (SEQ ID NO:97);

AATTCTCATTAGAGCT (SEQ ID NO:98);

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ECORI SSEI Saci

3'-TCGAGATTACTCTTAA-5' (SEQ ID NO:100) 5'-CTAATGAG-3' (SEQ ID NO:99)

manufactured by Stratagene) which had been digested in electrophoresis to recover a fragment of about 600 bp in size which encodes amino acid residues 1 to 174 of human TPO, and he fragment was subsequently purified using Prep-A-Gene advance with Xbal and EcoRI (E. coli DH5a was used as the In the same manner, the clone pBL(XS)(1-174) was digested with Xbal and Hindlll to recover a fragment of about 550 bp in size which encodes amino acid residues 1 to 174 of human A-136490) which had been digested in advance with Xbal and Hindlil (E. coli DH5 α was used as the host). A clone obtained in digested with Xbal and EcoRI and subjected to 2% agarose gel TPO, and the fragment was subsequently purified using Prep-The clone pUC18(XS)(1-174) obtained above was A-Gene DNA Purification Kit and sub-cloned into pCFM536 (EP-ONA Purification Kit and sub-cloned into pBluescript IISK+ host). A clone obtained in this way was named pBL(XS)(1-174) this way was named pCFM536/hT(1-174).

PCR reaction (incubation at 96 c for 2 minutes, repetition of a incubations at 95 c for 1 minute, at 41 c for 1 minute and at was carried out using pCFM536/hT(1-174) as a template and we PCR primers GEZ1 and GEX3. The human TPO encoding ragment obtained in this way was digested with Nael and total of 22 cycles of reaction, each cycle consisting of 72.c for 1 minute, and final incubation at 72.c for 7 minutes) The following experiment was carried out in order 2T (manufactured by Pharmacia) which is an expression vector to effect expression of GST-TPO(1-174) making use of pGEXof glutathione-S-transferase (GST) fusion protein.

then a clone, named pGEX-2T/hT(1-174), which encodes recover a fragment of about 550 bp in size, the fragment was subsequently purified using Prep-A-Gene DNA Purification Kit nucleotide sequence analysis to prepare a transformant for use in the expression of GST-TPO(1-174). Oligonucleotide sequences of the primers used in the PCR reaction are as and cloned into pGEX-2T which has been digested in advance with EcoRI and Smal (E. coli DH5 was used as the host) and correct human TPO nucleotide sequence was selected by EcoRI and subjected to 2% agarose gel electrophoresis follows:

GEX1: 5'-ATC GCC GGC TCC GCC AGC TTG TGA C-3' (SEQ ID NO:101) (prepared by adding a Nael recognition sequence to the sequence of positions 21 to 39 in SEQ ID NO:10); and

(SEQ ID NO:102) (an antisense primer corresponding to the GEX3: 5-GCC GAA TTC TCA TTA GAG CTC GTT CAG TGT-3' sequence of positions 523 to 549 in SEQ ID NO:10).

septide and a human TPO (amino acid residues 1 to 174). The DNA sequence which encodes thrombin recognition peptide and This expression plasmid contains a DNA sequence which encodes GST protein followed by a thrombin recognition human TPO (amino acid residues 1 to 174) are shown in the Sequence Listing (SEQ ID NO:10).

<Example 40>

Expression of GST-TPO(1-174) in E. coli

cultured overnight at 37°C on a shaker using 60 ml of LB medium containing 50 µg/ml of ampicillin, and 25 ml portion of the resulting culture broth was added to 1,000 ml of LB medium containing 50 µg/ml of ampicillin and cultured at 37 c on the shaker until OD at Agog reached 0.7 to 0.8. The transformant prepared in Example 39 was Thereafter, IPTG was added to the culture broth to a final concentration of 0.1 mM and the shaking culture was continued - 184

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for additional 3 hours in order to induce expression of GST-TPO(1-174).

<Example 41> Purification of GST-TPO(1-174) expressed in E. coli. and confirmation of its biological activity.

A 5.9 g portion of frozen cells of the recombinant strain which produces GST-TPO(1-174) derived from human TPO nucleotide sequence-encoding clone pGEX-27/hT(1-174) were suspended in 10 ml of water and disruptured using a high pressure disintegrator. By subjecting the suspension to centrifugation, GST-TPO(1-174) was recovered in the precipitated fraction and most parts of contaminated proteins, cell components and the like were removed. Next, the thus recovered precipitated fraction containing GST-TPO(1-174) was suspended in 5 ml of water to which were subsequently added, with stirring, 6 ml of 1 M Tris buffer (pH 8.5), 120 ml of 10 M urea and 16 ml of water. After 5 minutes of stirring to solubilize the contents, the resulting solution was evenly divided into 4 portions to carry out the following steps (1) to (4).

(1) One of the divided samples was diluted by a factor of 10 with 20 mM Tris buffer having a pH value of 8.5. To this were added reduced type glutathione and oxidized type glutathione to respective final concentrations of 5 mM and 0.5 mM, followed by overnight incubation at 4 ° ° c. The thus prepared mixture was subjected to centrifugation to recover GST-TPO(1-174) as the supernatant fluid which was subsequently diluted by a factor of 2 with 20 mM sodium citrate buffer having a pH value of 5.5 and then adjusted to pH 5.5 with acetic acid. GST-TPO(1-174) in the solution was applied to SP Sepharose Fast Flow (manufactured by Pharmacia Biotech, catalog No. 17-0729-01) cation exchange column which has been equilibrated in advance with 20 mM sodium

sodium citrate buffer pH 5.5, elution of GST-TPO(1-174) was effected using 20 mM sodium citrate buffer pH 5.5 containing 500 mM sodium chloride. A 129 ml portion of the eluate was mixed with 2.6 ml of 1 M Tris buffer pH 8.5, and the resulting nixture having a pH value of 8.1 was applied to Glutathione The thus recovered non-adsorbed fraction was diluted by a applied to SP Sepharose Fast Flow cation exchange column Sepharose 4B (manufactured by Pharmacia Biotech, catalog No. 17-0756-01) column to adsorb GST-TPO(1-174). After washing the column with PBS, elution of GST-TPO(1-174) was effected using 20 mM Tris buffer pH 8.5 containing 10 mM of educed type glutathione. The resulting eluate was mixed with 37 NIH units of thrombin, allowed to stand for 4 hours at room emperature, diluted with PBS by a factor of 10 and then applied to Glutathione Sepharose 4B column to adsorb digested actor of 3 with 20 mM sodium citrate buffer pH 5.5 and which has been equilibrated in advance with the same buffer, and the compound of interest was eluted with a linear density gradient of 0 to 500 mM sodium chloride dissolved in the same GST and to recover TPO(1-174) in the non-adsorbed fraction. citrate buffer pH 5.5. After washing the column with 20

(2) All operations of the step (1) were carried out in the presence of 0.1% polysorbate 80.

(3) One of the divided samples was diluted by a factor of 10 with 20 mM Tris buffer having a pH value of 8.5. To this were added reduced type glutathione and oxidized type glutathione to respective final concentrations of 5 mM and 0.5 mM, followed by overnight standing at 4 ° c. The thus prepared mixture was subjected to centrifugation to recover GST-TPO(1-174) as the supernatant fluid which was subsequently diluted by a factor of 2 with 20 mM sodium citrate buffer having a pH value of 5.5 and then adjusted to pH 5.5 with acetic acid. GST-TPO(1-174) in the solution was applied to SP

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was subsequently eluted with a linear density gradient of 0 to 5.5. After washing the resin with 20 mM sodium citrate buffer sodium citrate buffer pH 5.5 containing 500 mM sodium chloride. The eluate was adjusted to pH 8 with 1 M Tris buffer peptide linker, allowed to stand for 4 hours at room temperature, diluted by a factor of 5 with 20 mM sodium citrate buffer pH 5.5 and then applied to SP Sepharose Fast Flow cation exchange column which has been equilibrated in advance with the same buffer, and the compound of interest Sepharose Fast Flow cation exchange resin which has been equilibrated in advance with 20 mM sodium citrate buffer pH pH 5.5, elution of GST-TPO(1-174) was effected using 20 mM 5H 8.5, mixed with 320 NIH units of thrombin to remove GST polypeptide sequences via cleavage at a thrombin recognition 500 mM sodium chloride sissolved in the same buffer.

(4) All operations of the step (3) were carried out in the presence of 0.1% polysorbate 80.

sequence to be expressed as the pGEX-2T/hT(1-174)-derived The deduced amino acid sequence of the esulting TPO was [Gly-1]TPO given the sequence of the (purity, 1 to 20%) as one of the main protein bands of the carried out by subjecting it to SDS-PAGE and PVDF membrane ransfer in accordance with the procedure described in Example 1, it was confirmed that this band contains a TPO hrombin recognition peptide linker and the known activity of Each of the fractions eluted at sodium chloride densities of about 200 to 400 mM by the SP Sepharose Fast Flow cation exchange column chromatography was thoroughly dialyzed against IMDM culture medium and then evaluated by he rat CFU-MK assay system. When the fraction was analyzed by SDS-PAGE in the presence of a reducing agent, a band corresponding to a molecular weight of 19 kd was detected raction. When N-terminal sequence analysis of this band was hrombin on the linker. fusion protein:

Construction of vector for use in E. coli expression of human TPO (amino acid residues 1 to 163) having substitutions in the position 1 (Ser -> Ala) and position 3 (Ala -> Val) Example> 42

gel electrophoresis to recover a fragment of about 1,000 bp in SK+ (manufactured by Stratagene) which has been digested in recognition sequence to position 163 amino acid (positions 366 to 489) was changed to E. coli dominant codons. Synthetic oligonucleotides 13 to 20 shown below were prepared, and each pair of synthetic oligonucleotides 13 and 14, 15 and 16 a solution composed of 0.1 mM ATP, 10 mM Tris-acetate, 10 7.5), 100 mM MgCl₂ and 500 mM NaCl, the resulting solution was heated in a boiling water bath for 3 minutes and then allowed to cool to room temperature to obtain a doubledouble-stranded DNA fragments consisting of 13 and 14, 15 and 16 and 17 and 18 were ligated using DNA Ligation Kit manufactured by Takara Shuzo), and PCR was carried out using he thus ligated product as a template and the synthetic oligonucleotides 19 and 20 as primers. The thus obtained PCR product was digested with BamHI and HindIII and subjected to The clone pUC18(XE)(1-332) prepared in Example 39 was digested with Xbal and EcoRI and subjected to 2% agarose size which encodes amino acid residues 1 to 332 of human TPO, and the fragment was subsequently purified using Prep-A-Gene DNA Purification Kit and sub-cloned into pBluescript II advance with Xbal and EcoRI (E. coli DH5 α was used as the Next, a region of the clone pBL(XE)(1-332) from its BamHI and 17 and 18 were phosphorylated in the same tube containing mM Mg-acetate and 50 mM K-acetate. After further addition of 1/10 volume of a solution composed of 100 mM Tris/HCI (pH stranded DNA fragment. Next, the thus obtained three pairs of host). A clone obtained in this way was named pBL(XE)(1-332).

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130 bp in size using Prep-A-Gene DNA Purification Kit. This was sub-cloned into pBL(XE)(1-332) which has been digested in advance with BamHI and HindIII (E. coll DH5 was used as the 2% agarose gel electrophoresis to recover a fragment of about host). A clone having the following nucleotide sequences was selected from the thus obtained clones by sequence analysis and named pBL(XH)(1-163):

3: 5-GATCCGAACGCTATCTTCCTGTCTTCCAGCACCTGCTGCGT-3

(SEQ ID NO:103);

(SEQ ID NO:104);

14: 5-TTTGCCACGCAGCAGGTGCTGGAAAGACAGGAAGATAGCGTTCG3

15: 5-GGCAAAGTTOGTTTCCTGATGCTGGTTGGCGGTTCTACCCTG-3"

(SEQ ID NO:105);

16. 5-ADGCACAGGGTAGAAQQGQCAAQCAGCATCAGGAAAQGAACG

17: 5-TGOGTTOGTOGGGGGGGCGCACCACTGCTGTTCOGTCTTAATGAA:3

(SEQ ID NO:106);

(SEQ ID NO:107);

IR 5-AGCTTTCATTAAGAOGGAACAGCAGTGGTTGGOGGOGOCOCGACGA3

(SEQ ID NO:108);

19: 5-AAGGATCCGAACGCTATCTTCCTG-3" (SEQ ID NO:109); and 20: 5-AGAAGCTTTCATTAAGACGGAACA-3' (SEQ ID NO:110).

N-terminal by protease, an expression vector was constructed of human TPO (positions 1 to 163) and preventing hydrolysis of or use in the expression of a mutation type human TPO positions 1 to 163) (to be referred to as "h6T(1-163)" [[Ala1, Val3]TPO(1-163)) and, at the same time, encoding Lys Ala¹, Val³]TPO(1-163)). Four synthetic oligonucleotides For the purpose of increasing expression quantity hereinafter) having substitutions at the position 1 (Ser to Ala) and position 3 (Ala to Val) of human TPO (positions 1 to 163) at the -1 position and Met at the -2 position ([Met-2, Lys-1, shown below were prepared, and synthetic oligonucleotides 2-9 and 3-3 were phosphorylated with T4 kinase (manufactured

ragments 1-9 and 2-9 and 3-3 and 4-3 were annealed in a order to make these synthetic oligonucleotides into doublestranded DNA fragments, each pair of the single-stranded DNA solution composed of 10 mM Tris/HCI (pH 7.5), 10 mM MgCl₂ reaction was sub-cloned into pBL(XH)(1-163) which has been substituted by the following synthetic oligonucleotide was by Pharmacia) in a solution composed of 1 mM ATP, 10 mM and 50 mM NaCl. Next, the thus obtained two pairs of doublestranded DNA fragments consisting of 1-9 and 2-9 and 3-3 and 4-3 were treated with DNA Ligation Kit (manufactured by Takara Shuzo). The DNA fragment thus obtained by the ligation digested in advance with Xbal and Nrul, and a clone correctly selected by sequence analysis (E. coli DH5 α was used as the Tris-acetate, 10 mM Mg-acetate and 50 mM K-acetate. nost) and named pBL(XH)h6T(1-163):

1-9: 5'-CTAGAAAAACCAAGGAGGTAATAAATAATGAAAGCACCTGTACCA-3'

(SEQ ID NO:112);

2-9: 5 -caggregracaggrecritcattartrattaccrcctrggrtttt-3

3-3: 5'-ccrccargreattracescrccrsrcraaacrecrece-3'

(SEQ ID NO:113); 4-3: 5'-cccaccagtttagacaggacccgtaaatcacatg-3'

(SEQ ID NO:114);

1 2

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CTAGAAAAA CCAAGGAGGT AATAAATAAT GAAAGCACCT TITITI GGTTCCTCCA TIAITIATIA CITTCGTGGA

57 Xbal 2 귀 8

GTACCACCTG CATGTGATTT ACGGSTCCTG TCTAAACTGC TGCG CATGGTGGAC GTACACTAAA TGCCCAGGAC AGATTTGACG ACGC

-1

(SEQ ID NO:115)

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E. coli. and confirmation of its biological activity

resulting solution was diluted 10 times with 20 mM Tris of stirring at room temperature to solubilize the contents, the fractions was analyzed by SDS-PAGE in the presence of a this band was carried out by subjecting it to SDS-PAGE and in Example 1, it was confirmed that this band contains a TPO SFU-MK assay system, TPO activity was found in all fractions raction was recovered and most parts of contaminated proteins, cell components and the like were removed. The thus ecovered precipitated fraction containing h6T(1-163) was urther addition of urea (final concentration, 8 M), guanidine sarcosinate (final concentration, 2%). After 5 to 20 minutes buffer having a pH value of 8.5 and subjected to protein glutathione and copper sulfate were used as additives in addition to air oxidation. By centrifugation, h6T(1-163) was recovered in the supernatant fluid. When each of the recovered educing agent, a band corresponding to a molecular weight of 18 kd was detected (purity, 30 to 40%) as the main protein band of each fraction. When N-terminal sequence analysis of PVDF membrane transfer according to the procedure described dialyzed against IMDM culture medium and evaluated by the rat A 3.6 g portion of frozen cells of the h6T(1-163) oroducing recombinant strain were suspended in 10 ml of water and disruptured using a high pressure disintegrator. By subjecting the suspension to centrifugation, the precipitated suspended in 7 ml of water, and 3 ml of 1 M Tris buffer (pH 8.5) was added to the suspension with stirring, followed by nydrochloride (final concentration, 6 M) and sodium N-lauroyl refolding by overnight oncubation at 4 c. In this case, sequence to be expressed as the pCMF536/h6T(1-163)-derived mutation type TPO. When each fraction was thoroughly

Purification of h6T(1-163) expressed in

Example 44>

n a dose-dependent fashion.

encodes amino acid residues 1 to 163 of the mutation type human TPO, and the fragment was subsequently purified using and Hindill (E. coli JM109 which has been transformed in and an E. coli strain having this expression vector was used as a transformant for use in the expression of the mutation type human TPO, namely h6T(1-163). This expression plasmid and Hindlil to recover a fragment of about 500 bp in size which Prep-A-Gene DNA Purification Kit and cloned into pCFM536 (EP-A-136490) which has been digested in advance with Xbal advance with pMW1 (ATCC No. 39933) was used as the host). A clone obtained in this way was named pCFM536/h6T(1-163), contains the DNA sequence shown in the Sequence Listing (SEQ The clone pBL(XH)(1-163) was digested with Xbal

Expression of h6T(1-163) in E. coli

OD at A₆₀₀ reached 1.0 to 1.2. Thereafter, about 330 ml of LB . Expression of the expression plasmid pCFM536 is controlled by APL promoter which by itself is under control of cl857 repressor gene. The transformant obtained in < Example 42> was cultured overnight at 30 c on a shaker using 60 ml of LB medium containing 50 µg/ml of ampicillin and 12.5 µg/ml of tetracycline, and 25 µl portion of the resulting culture broth was added to 1,000 ml of LB medium containing 50 ug/ml of ampicillin and cultured at 37.c on the shaker until nedium heated to 65°c was added to the culture broth to adjust the final medium temperature to 42.c, and the shaking culture was continued for 3 hours at 42.c to induce expression of h6T(1-163)

<Example 43>

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and detection of TPO by SDS-PAGE and western analysis Preparation of anti-TPO peptide antibody

means. In consequence, three regions which appear to be relatively useful as antigens (see below) were selected from multiple antigen peptide (MAP) type peptide was synthesized USA, 85, 5409 - 5413, 1988) using each of the selected regions, and then each of 2 rabbits was immunized 8 times with 100 µg of each of the thus synthesized peptides to obtain will render possible detection of TPO protein by immunological he confirmed TPO amino acid sequences, a quadruple-stranded Success in preparing anti-TPO peptide antibody n accordance with the procedure of Tam (Proc. Natl. Acad. Sci. respective antiserum samples.

Amino acid sequences contained in the synthesized peptide antigens

PRILINKILRDSYLLHBRLSQ (SEQ ID NO:116) (a) Rat IPO(9-28) (Peptide RT1 region)

(B. at Position 24 is originally S in the amino acid residues determined from the gene.)

(b) Rat TPO(46-66) (Peptide RT2 region)

FSLGEWKTQTEQSKAQDILGA (SEQ ID NO:117)

Rat TPO(163-180) (Peptide RT4 region) SRTSQLLTLNKFPNRLLD (SEQ ID NO:118) O

(LLD at Positions 178-180 is originally TSG in the amino residues determined from the gene.) ğcid

and anti-RT2 peptide were firstly subjected to a protein A ng and 1,920 mg, respectively), and 54 mg and 32 mg portions 8427) column chromatography to prepare IgG fractions (2,344 of the respective fractions were biotinylated by their coupling with active type biotin (NHS-LC-Biotin II; manufactured by Next, of these antiserum samples, anti-RT1 peptide (PROSEP-A; manufactured by Bioprocessing Ltd., catalog No.

on PVDF or nitrocellulose membrane in the same manner as PIERCE, catalog No. 21336). A recombinant TPO-containing sample was subjected to SDS-PAGE and then to electroblotting described in Example 1, and western analysis was carried out in the usual way using these biotinylated antibodies as first antibodies.

That is, the membrane after blotting was washed with a blocking agent (BlockAce; manufactured by Dainippon with a solution composed of 20 mM Tris-HCI and 0.5 M NaCI (pH Pharmaceutical, catalog No. uk-B25) for 60 minutes. Next, the membrane was treated for 60 minutes with TTBS solution containing 10 µg/ml of biotinylated anti-TPO peptide antibody, 0.05% BSA and 10% BlockAce and then washed twice with TTBS or 5 minutes each. Thereafter, the membrane was treated for 30 minutes with a solution prepared by diluting alkaline phosphatase-labeled avidin (manufactured by Leinco fechnologies, catalog No. A108) by a factor of 5,000 with ITBS solution containing 10% BlockAce and washed twice with TTBS for 5 minutes each and with TBS for 5 minutes, and then color development was effected by the use of an alkaline 70-6432). The western analysis described above was carried 7.5) (TBS) for 5 minutes and twice with 0.1% Tween 20containing TBS (TTBS) for 5 minutes each and then treated chosphatase substrate (manufactured by Bio-Rad, catalog No. out at room temperature.

hereof and h6T(1-163) which is a mutation type TPO As the results, it was able to recognize not only various recombinant rat TPO proteins expressed in COS 1 cells COS 1 cells and E. coli cells (illustratively, plasmid pHTP1- or plasmid pHTF1-derived human TPO expressed in COS 1 cells and N-glycosidic linkage-cleaved product thereof, GST-TPO(1-174) expressed in E. coli and thrombin-digested product but also various recombinant human TPO proteins expressed in expressed in E. coli, which have been described in Examples).



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In addition to the above, possibility for the preparation of anti-human TPO peptide antibodies was confirmed. Antibodies obtained by these techniques can be applied not only to western analysis but also to purification of TPO by antibody columns and every usually used antibody-aided immunological means.

Six regions in the human TPO amino acid sequence shown in SEQ ID NO:7, which were expected to have relatively suitable antigenicity, were selected (see Table 4). Tetrameric Multiple Antigen Peptide (MAP) type peptides, which correspond to the regions, respectively, were synthesized. Each peptide was immunized to two rabbits 8 times in an amount of 100 mg/time.

Separately from this, a monomeric peptide in which a cysteine residue had been bonded to the C-terminus of each peptide region shown in Table 4 was synthesized too. This peptide was then used as a test antigen to determine an antibody titer using enzyme immunoassay. As a result, the increase of the antibody titer was confirmed for the above prepared sera, so these sera were employed as antisera.

The antibodies were prepared by immunizing each antigenic peptide to two rabbits to produce antisera directed against the peptide. The resultant two anti-HT1 peptide antibodies are distinguishably referred to as anti-HT1-1 peptide antibody and anti-HT1-2 peptide antibody depending on the rabbit individuals.

An example of the purification of the anti-HT1-1 peptide antibodies will be illustrated below.

First, 30 mg of a monomeric peptide of HT1 to which has been bonded a cystein residue was coupled to 12 ml of Sulfo-Link coupling gel (Pierce, Catalogue No. 44895). Briefly, a peptide solution containing the antigen was coupled

for 15 minutes to the gel equilibrated with a coupling buffer (50 mM Tris, 5 mM EDTA-Na pH 8.5) of 6 volumes relative to the gel volume. Then, after incubation for 30 minutes, the gel was washed with the coupling buffer of 3 volumes relative to the gel volume. The coupling buffer containing 0.05 M L-cystein-HCl was added to the gel in an amount of 1 ml/ml gel to block unreacted radicals over 15 minutes. After incubation for 30 minutes, the gel was washed with the coupling buffer of 8 volumes relative to the gel volume. All the coupling reactions were carried out at room temperature. Thus the peptide was covalently bonded to the gel with a coupling efficiency of 28.3%, and then an antigen column with 0.8 mg of the peptide/ml gel was prepared.

76.7 ml (protein content 3620 mg) of the antiserum containing the anti-HT1-1 peptide antibody which had been bled from one of the rabbits with a total amount of 78.4 ml, was applied to the antigen column pre-equilibrated with 50 mM phosphate buffer (pH 8) containing 150 mM NaCl and 0.05% sodium azid, and then washed with the same buffer to afford 105.9 ml of a flow-through fraction (protein content 3680 mg). Then the adsorbed fraction was eluted with 0.1 M citrate buffer (pH 3.0) and immediately neutralized with 21.1 ml of 0.1 M carbonate buffer (pH 9.9), followed by concentration by means of ultrafiltration (using Amicon YM 30 membran) to give the purified anti-HT1-1 peptide antibody in a solution of 50 mM phosphate buffer (pH 8) containing 150 mM NaCl and 0.05% NaN₂.

Similarly, the following antibodies were prepared: anti-HT1-2 peptide antibody (60.0 mg); anti-HT2-1 peptide antibody (18.8 mg); anti-HT2-2 peptide antibody (8.2 mg); and the like. Also, anti-HT3 to -HT6 peptide antibodies could be prepared in the similar manner.

Three mg of affinity-purified anti-HT1-1 peptide antibody, anti-HT1-2 peptide antibody, anti-HT2-1 peptide

antibody or anti-HT2-2 peptide antibody was biotinylated by coupling to an activated blotin (Pierce, NHS-LC-Biotin II, Catalogue No. 21336).

All the above purified antibodies were found to recognize and detect human TPO by Western blotting (on nitrocellulose filter or PVDF) following SDS-PAGE of a recombinant human TPO standard partially purified from the culture supernatant of CHO cells wherein a gene encoding the amino acid sequence shown in Table 4 had been introduced and expressed.

Table 4 Human TPO amino acid sequences included in the

ruman TPV amino acid sequences included in the retrameric MAP type, synthetic peptide antigen

Human TPO (amino acid numbers 8-28) peptide HT1 region: DLRVLSKLLRDSHVLHSRLSQ (SEQ ID NO:119)

Human TPO (amino acid numbers 47-62) peptide HT2 region: SLGEWATCMEETKAQD (SEQ ID NO:120)

Human TPO (amino acid numbers 108-126) peptide HT3 region: LGTQLPPQGRTTAHKDPNA (SEQ ID NO:121) Human TPO (amino acid numbers 172-190) peptide HT4 region: NELPNRTSGLLETNFTASA (SEQ ID NO:122)

Human TPO (amino acid numbers 262-284) peptide HT5 region: SLPPNLQPGYSPSPTHPPTGQYT (SEQ ID NO:123) Human TPO (amino acid numbers 306-332) peptide HT6 region: PSAPTPTPTSPLLNTSYTHSQNLSQEG (SEQ ID NO:124)

Example 46>

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Preparation of anti-TPO peptide antibody column

Since the anti-rat TPO peptide antibody obtained in peptide antibody and anti-RT2 peptide antibody were linked to a chemically activated gel support in the following manner to prepare anti-TPO peptide antibody columns. Each of the two antibodies used as the materials was prepared from sera of Example 45 was able to recognize rat and human TPO molecules, immunoglobulin (IgG) fractions of the anti-RT1 peptide antibody and anti-RT1-2 peptide antibody, and the anti-RT2 peptide antibody prepared from other two rabbits RT2-2 peptide antibody. Since these antibodies were separately used for the antibody column preparation, a total of column" and "anti-RT1-2 antibody column" hereinafter), two antibody column) which was prepared by mixing anti-RT1-2 wo rabbits separately. That is, the anti-RT1 peptide antibody prepared from two rabbits was respectively named anti-RT1-1 was respectively named anti-RT2-1 peptide antibody and anti-5 columns were obtained, namely two anti-RT1 peptide antibody columns (to be referred to as "anti-RT1-1 antibody anti-RT2 peptide antibody columns (to be referred to as "anti-RT2-1 antibody column" and "anti-RT2-2 antibody column" hereinafter) and one antibody column (anti-RT1-2 + 2-1 mix peptide antibody with anti-RT2-1 peptide antibody and coupling the mixture with a gel.

Each antibody was dissolved in a solution of 50 mM Na phosphate and 0.15 M NaCl (pH 8.0) to a final concentration of 5 mg/ml (or 2.5 mg/ml each of evenly mixed anti-RT1-2 peptide antibody and anti-RT2-1 peptide antibody in the case of the anti-RT1 + 2 mix antibody column), and a 2.31 ml portion of the resulting solution was mixed with 1.54 ml in volume of a swelled formyl-activated gel (Formyl-Cellulofine, manufactured by Chisso) and subjected to 2 hours of coupling reaction at 4 · c. To this was added 1.1 ml of a 10 mg/ml solution of a reducing agent (trimethylamine borane (TMAB);

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The M-07e assay system was used as the in vitro

followed by 6 hours of additional coupling reaction and subsequent centrifugation to recover the gel portion. A 10, ml was recovered again by centrifugation, and unreacted antibody the resulting gel was mixed with 4.6 ml of a blocking solution (0.2 M Na phosphate and 1 M ethanol amine, pH 7.0) and 1.1 ml of the reducing agent solution and treated at 4.c for at least 2 Thereafter, the gel was washed with purified water and DPBS portion of purified water was added to the gel portion which molecules were removed by repeating this step 4 times. Next, using a centrifuge, packed in a small column tube, washed with a 3 M potassium thiocyanate solution and a 0.1 M glycine-HCl hours to perform blocking of active groups of unreacted gel. manufactured by Seikagaku Kogyo, catalog No. 680246) (pH 2.5) solution, equilibrated with DPBS and then stored.

RT1-2 + 2-1 mix antibody column, the coupling efficiency of consequence, since it was confirmed that the coupling depending on the antibody origins and difference in the intigens, the experiments shown in the following Example 47 respective . IgG fractions reached 97.4%, 95.4%, 98.4%, 98.3% and 99.4%. In addition, amounts of IgG fractions coupled per gel volume were 5.6 mg/ml gel, 5.8 mg/ml gel, 5.7 mg/ml gel, efficiency and coupling quantity do not significantly vary in the 5 anti-TPO peptide antibody gels of the anti-RT1-1 antibody column, anti-RT1-2 antibody column, anti-RT2-1 antibody column, anti-RT2-2 antibody column and anti-5.7 mg/ml gel and 2.9 mg/ml gel, respectively. vere carried out making use of these antibody columns.

Example 47>

using anti-TPO antibody column and then by reverse phase column chromatography, and confirmation of its biological Purification of TPO derived from culture supernatant obtained by transfecting COS 1 cells with expression vector pHTP1. activity

Partially purified samples of TPO derived from the culture supernatant obtained in Example 35 by transfected COS 1 cells main TPO fraction, namely a passed-through fraction F1 and a ultrafiltration unit (manufactured by Amicon) to obtain a assay for the evaluation of the following purification samples. with the expression vector pHTP1 (fractions other than the 5H 5.8, of Macro-Prep Methyl HIC column and fractions F1 and F3 of SP Sepharose Fast Flow column) were combined to prepare a TPO subpool fraction (5,463.79 ml; protein concentration, 0.490 mg/ml; total protein, 2,676 mg; relative (manufactured by Amicon) to make it finally into 120.2 ml of a DPBS solution containing 0.05% sodium azide. Next, all of the 5 anti-TPO peptide antibody gels prepared in Example 46 were mixed and made into a single antibody column (1.6 cm in diameter and 4.8 cm in bed height, to be referred to as anti-RT1 + 2 mix antibody column), and the TPO subpool fraction After completion of the sample loading, elution was carried out with DPBS to collect eluates until UV absorption became sufficiently small, and the thus collected eluates were concentrated using the YM-10 membrane-equipped bassed-through fraction F1 (82.62 ml; protein concentration, raction F3 eluted after F2 by washing with 20 mM Na citrate, activity, 12,100; and total activity, 32,380,000) and concentrated using an ultrafiltration unit (Omega Ultraset, nominal molecular weight cutoff of 8,000, manufactured by Filtron), solvent of the concentrated fraction was exchanged with DPBS and then the small-volumed sample was treated with an YM-10 membrane-equipped ultrafiltration unit was applied to the column at a flow rate of 0.033 ml/min. 14.3 mg/ml; total protein, 1,184 mg; relative activity, 67,500; and total activity, 79,900,000). Next, a column-adsorbed raction F2 (92.97 ml; protein concentration, 0.12 mg/ml; total protein, 11.1 mg; relative activity, 257,100; and total activity,

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increase in the relative activity. That is, using 0.1% TFA as a 2,860,000) was eluted with an acidic eluting solution (0.1 M TPO was further purified because it showed about 20 times developing solvent A and 1-propanol containing 0.05% TFA as a antibody column with 1/10 volume of the developing solvent B and applying the mixture at a flow rate of 0.4 ml/min to the Capcell Pak Cl 300A column which has been equilibrated in 50 minutes with a linear density gradient of from 20% B to glycine-HCl, pH 2.5). Though the passed-through fraction F1 still contained a TPO activity, the antibody column-adsorbed developing solvent B, Capcell Pak CI 300A (manufactured by Shiseido, catalog No. Cl TYPE:SG300A; 4.6 mm in diameter and 150 mm in bed height, connected with a precolumn of 4.6 mm was carried out, by mixing the fraction F2 obtained from the advance with 20% B. After completion of the sample loading, elution was carried out for 5 minutes with 20% B and then for 40% B, and the eluates were collected in polypropylene tubes n diameter and 35 mm in bed height) column chromatography in 1 ml (2.5 min) portions.

A 2 µl portion (1/500 fraction) of the eluate in ultrafiltration to obtain 0.25 ml of IMDM assay culture solution containing 0.02% HSA for use in the identification of TPO active fractions by the assay. As the result, strong TPO analysis in accordance with the procedure described in Example 45>, the presence of TPO was confirmed which had an under a reduced condition based on DPC III molecular weight markers, as well as the presence of other molecules having each tube was mixed with HSA and concentrated by activity was found in samples of tube Nos. 20 to 23 (within the addition, when these samples were subjected to western apparent molecular weight of 60,000 to 70,000 when measured respective molecular weights of 32,000 to 43,000 and 20,000 range of 28.5 and 32.5% as propanol concentration).

Example 48>

of biological activity of TPO derived from culture supernatant obtained by transfecting COS 1 cells with expression vector pHTP1 and purified up to the step of Capcell Pak Cl 300A column Confirmation

administered in the same manner. Just before the by Toa of the administration in comparison with the value prior to the (2.6 ml). In order to examine biological TPO activity of this sample, in vivo assay was carried out. That is, the active weeks of age), each group including 4 animals, daily for 5 days at a dose of 100 µl (having a total activity of 110,000 blood was collected from the eyeground to measure platelet administration and was 1.23 times higher than the case of the namely tube Nos. 20 to 23 (within the range of 28.5 and 32.5% as propanol concentration) of Capcell Pak Cl 300A column, were pooled, mixed with 0.21 ml of glycerol and then to a volume of 1 ml with DPBS, buffer-exchanged with DPBS 0.01% HSA, thereby finally obtaining a TPO active fraction FA fraction was subcutaneously administered to ICR male mice (8 yo Denshi). In the TPO-treated group, platelet counts ncreased by a factor of about 1.42 in average after completion control group with a significant difference (p <0.05, Student t-On the basis of these results, it was The TPO active fractions purified in Example 36, concentrated by centrifugation evaporation. This was mixed with 0.21 ml of 6 M guanidine hydrochloride solution, diluted solution containing 0.01% HSA by Sephadex G25 column (NAP-10, manufactured by Pharmacia Biotech, catalog No. 17-0854-01) and then mixed with 1.1 ml of DPBS solution containing measured by M-07e assay system) per animal. As a control, 100 µl of DPBS containing 0.01% of HSA was subcutaneously administration and on the next day of the final administration, counts using a microcell counter (F800, manufactured est) between them.

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mammalian cells has the ability to increase platelet counts in confirmed that the aforementioned human TPO produced by

<Example 49>

Confirmation of biological activity of crude TPO fraction derived from 33 liters of culture supernatant obtained by transfecting COS 1 cells with expression vector pHTP1 and partially purified by cation exchange column

(1) The M-07e assay system was used as the in samples. In the same manner as described in Example 36, a total of 33 liters of serum-free culture supernatant was obtained by transfecting COS 1 cells with the expression in molecular weight indicates a possibility that the culture vector pHTP1 and filtered through a 0.22 µm filter to recover the supernatant. This was concentrated by a factor of about Millipore) and mixed with about 1 mM in final concentration of a protease inhibitor p-APMSF to obtain a sample of 2,018 ml in 429,100,000). Next, this was concentrated repeatedly using an cutoff of 30,000; manufactured by Filtron), thereby obtaining a vitro assay for the evaluation of the following purification 10 with an ultrafiltration unit (PLGC pellicon Cassette, nominal molecular weight cutoff of 10,000, manufactured by volume (protein concentration, 3.22 mg/ml; total protein, 6,502 mg; relative activity, 66,000; total activity, ultrafiltration unit (Omega Ultraset, nominal molecular weight fraction of 30,000 or more in molecular weight (volume, 1,190 ml; protein concentration, 2.54 mg/ml; total protein, 3,020 ng; relative activity, 82,500; total activity, 249,000,000) and a fraction of 30,000 or less in molecular weight (volume, 2,947 ml; protein concentration, 0.471 mg/ml; total protein, The presence of TPO activity in the fraction of 30,000 or less supernatant contains TPO whose molecular weight has been 1,402 mg; relative activity, 4,500; total activity, 6,310,000)

reduced during its expression in or secretion from the animal cells. On the other hand, the fraction of 30,000 or more in molecular weight was buffer-exchanged with 20 mM sodium citrate buffer (pH 6.1) and applied at a flow rate of 5 ml/min to SP Sepharose Fast Flow (2.6 cm in diameter and 29 cm in 17-0729-01) column which has been equilibrated in advance with 20 mM sodium citrate buffer (pH 6.1). After completion of the sample loading, the column was washed and eluted with sodium citrate buffer (pH 6.1) as a developing solvent A and a solution composed of the developing solvent A and 1 M NaCl as a developing solvent B, elution was carried out at a flow rate of 3 ml/min with a linear gradient of from 0% B to 50% B for and the eluates were collected in polypropylene tubes in 30 ml (10 min) portions. When a portion of each of these eluates was the activity, elution of TPO activity was found within a broad range. In consequence, fractions eluted with a NaCl raction, were pooled as a fraction F1, and the main TPO Jolume of the fraction F1 was 1,951 ml (protein concentration, 2.05 mg/ml; total protein, 3,994 mg; relative activity, 13,500; total activity, 53,900,000) and that of F2 sed height, manufactured by Pharmacia Biotech, catalog No. 20 mM sodium citrate buffer (pH 6.1). Next, using 20 mM 215 minutes and then from 50% B to 100% B for 20 minutes, checked by the M-07e assay system to examine distribution of concentration of 50 mM or lower, including the passed-through was 649.8 ml (protein concentration, 1.11 mg/ml; total ractions eluted with 50 to 1,000 mM NaCl as a fraction F2. protein, 721 mg; relative activity, 268,000; total activity, 93,000,000).

SP Confirmation of biological activity of Sepharose Fast Flow fraction F2

A 2.5 ml portion of the SP Sepharose Fast Flow raction F2 separated in the above step (1) was bufferexchanged with 3.5 ml of DPBS solution containing 0.01% HSA

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of this sample (protein concentration, 1.46 mg/ml) was examined in an in vivo assay. That is, the active fraction was neasured by the M-07e assay system) per animal. As a subcutaneously administered in the same manner. Just before counts using a microcell counter (F800, manufactured by Toa each group including 4 animals, daily for 5 consecutive days at a dose of 100 µl (having a total activity of 123,000 control, 100 µl of DPBS containing 0.01% of HSA was yo Denshi). In the TPO-administered group, platelet counts of the administration in comparison with the value prior to the between them. These results demonstrated that the aforementioned human TPO produced by mammalian cells has by Sephadex G25 column (NAP-25, manufactured by Pharmacia Biotech, catalog No. 17-0852-01), and biological TPO activity subcutaneously administered to ICR male mice (8-weeks age), he administration and on the next day of final administration, blood was collected from the eyeground to measure platelet increased by a factor of about 1.29 in average after completion administration and were 1.12 times higher than in the control group with a significant difference (p <0.05, Student t-test) he ability to increase platelet counts.

cExample 50>

Construction of recombinant vector, pDEF202-ghTPO, for use in the expression of human TPO chromosomal DNA in CHO cells

A vector pDEF202 was digested with restriction enzymes Kpnl and Spel and then subjected to agarose gel electrophoresis to isolate a fragment containing mouse DHFR mini gene and SV40 polyadenylation signal, and the expression vector pDEF202-ghTPO was obtained by ligating, using T4 DNA ligase (manufactured by Takara Shuzo), the thus obtained fragment with a vector DNA which has been prepared by removing an SV40 polyadenylation signal-containing region

from a plasmid pEFHGTE by its treatment with restriction enzymes $K\rho nl$ and $S\rho el.$ This plasmid contains SV40 replication origin, human elongation factor 1- α promoter, SV40 early polyadenylation region for transcription of human TPO gene, mouse DHFR mini gene, pUC 18 replication origin and β -lactamase gene (Amp¹), and human TPO chromosomal DNA is connected to a downstream site of the human elongation factor 1- α promoter.

Example 51>

Expression of human TPO chromosomal DNA in CHO cells

CHO cells (a dhfr strain; Urlaub and Chasin, *Proc. Natl. Acad. Sci. USA*, vol.77, p.4216, 1980) were grown by culturing them in an α-minimum essential medium (α-MEM(-), supplemented with thymidine and hypoxanthine) containing 10% fetal bovine serum using a plate of 6 cm in diameter (manufactured by Falcon), and the resulting cells were subjected to transfection by means of a calcium phosphate method (CellPhect, manufactured by Pharmacia).

That is, 10 μg of the plasmid pDEF202-ghTPO prepared in Example 50 was mixed with 120 μ l of buffer A and 120 μ l of H₂O, and the resulting mixture was incubated at room temperature for 10 minutes. This solution was further mixed with 120 μ l of buffer B and incubated at room temperature for additional 30 minutes. The thus prepared DNA solution was put in the plate and cultured for 6 hours in a CO₂ incubator. After removing the medium and washing the resulting plate twice with a-MEM(-), a-MEM(-) containing 10% dimethyl suffoxide was added to the plate and treated for 2 minutes at room temperature. Next, non-selection medium (a-MEM(-) op. cit., supplemented with hypoxanthine and thymidine) containing 10% dialyzed fetal bovine serum was added and cultured for 2 days, and then selection was carried out using a

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selection medium (lpha-MEM(-), free from hypoxanthine and thymidine) containing 10% dialyzed fetal bovine serum. Selection was carried out by treating the cells with trypsin, dispensing the treated cells in one plate of 6 cm in diameter into 5 plates of 10 cm in diameter or 20 plates of 24 well plates and continuing culturing of the cells while changing the selection medium every 2 days. The presence of human TPO activity was confirmed in the culture supernatant in the plates or wells in which proliferation of the cells was observed, when the human TPO activity was measured by the Ba/F3 assay.

In this instance, transfection of CHO cells may also be effected by carrying out co-transfection of CHO cells with DEPHGTE and pMG1.

<Example 52>

of human TPO protein (amino acid residues of positions 1 to 163) (to be referred to as "hMKT(1-163)" hereinafter) in which a Lys residue and a Met residue were respectively added to the Construction of vector for use in the expression in E. coli 1 and -2 positions, and confirmation of its expression In order to effect expression of a protein whose 1human TPO protein (amino acid residues of positions 1 to 163), a vector pCFM536/hMKT(1-163), also referred to as [Met-2, Lys-1]TPO(1-163), was constructed for use in the expression of hMKT(1-163) in E. coli, in the same manner as described in Example 42 using the following synthetic oligonucleotides 1and 3-position amino acid residues are the same as those of 13, 2-13, 3-3 and 4-3:

1-13: 5'-CTAGARARAACCAAGGAGGTAATAAATAATGAAATCTCCTGCACCA-3'

(SEQ ID NO:125);

SEQ ID NO:126);

2-13: 5'-CAGGIGGIGCAGGAGATITCATIAITIATIACCICCTIGGITTITI-3'

5'-ccrecatetalgatitacegetoctetctaaactectece-3' 3-3:

(SEQ ID NO:127);

(SEQ ID NO:128); 5'-cgcagcagtttagacaggaccggtaaatcacatg-3

4-3:

6 1-13 20 TITITI GGIICCICCA ITAITIAITA CITTAGAGGA <u>1</u>

CTAGAAAAA CCAAGGAGGT AATAAATAAT GAAATCTCCT

ទ

TeqX

8 2 П e

GCACCACCTG CATGTGATTT ACGGGTCCTG TCTAAACTGC TGCG CGTGGTGGAC GTACACTAAA TGCCCAGGAC AGATTTGACG ACGC

(SEQ ID NO:129)

This expression plasmid contains a DNA sequence shown in the Sequence Listing (SEQ ID NO:12). Thereafter, expression of hMKT(1-163) was induced in the same manner as described in Example 43.

then subjected to N-terminal amino acid sequence analysis, it When the thus obtained expression protein was subjected to SDS-PAGE, transferred on a PVDF membrane and was confirmed that this protein contains the amino acid sequence of hMKT(1-163) to be expressed.

<Example 53>

Refolding of human TPO, h6T(1-163), derived from human TPO expressed in E. coli. using quanidine hydrochloride and olutathione, and purification and confirmation of its biological activity

producing recombinant strain prepared in Example 43 were suspended in 3 ml of water and disrupted using a high pressure A 1.2 g portion of frozen cells of the h6T(1-163)-

contaminated proteins, cell components and the like were

the precipitated fraction was recovered and most parts of

disintegrator. By subjecting the suspension to centrifugation

1 ml portion of 1 M Tris buffer (pH 8.5) was added to the

h6T(1-163) was suspended in 4 ml in final volume of water. A

removed. The thus recovered precipitated fraction containing

of 8 M guanidine hydrochloride and subsequent 5 minutes of stirring at room temperature to solubilize the contents. The resulting solution was diluted 10 times with 20 mM Tris buffer (pH 8.5), 5 mM of reduced type glutathione and 0.5 mM of oxidized type glutathione were dissolved in the diluted solution with stirring and then the thus prepared solution was incubated overnight at 4.c. By centrifugation, h6T(1-163) was recovered in the supernatant fluid. A 160 ml portion of the thus obtained supernatant fluid was concentrated using YM10 ultrafiltration membrane (manufactured by Amicon) and subjected to buffer exchange with DPBS to obtain 3.4 ml in

suspension with stirring, followed by further addition of 20 ml

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difference (p <0.001, Student t-test) between them. These esults demonstrated that the human TPO produced by E. coli actor of about 2.15 in average on the next day of the completion of the administration in comparison with the value orlor to the administration, and such an effect remained unchanged even after 3 days of the completion of the administration showing a 2.13 times higher value. Platelet counts on the next day of and 3 days after the completion of the administration was 1.73 and 1.80 times higher in TPOreated group than in the control group with a significant and prepared by the above method has the ability to increase he TPO-administered group, platelet counts increased by platelet counts in vivo.

Example 542

and purification and confirmation of its biological activity Refolding of human TPO. h6T(1-163) expressed in E. coli. using sodium N-lauroyl sarcosinate and copper sulfate,

11.25 µl of 1 M DTT, 38 µl of 0.5 M EDTA and 0.38 ml of 10% sodium deoxycholate solution were added to the suspension with stirring, and the resulting mixture was stirred at room This was then subjected to producing recombinant strain prepared in Example 43 were suspended in 3 ml of water, disrupted using a high pressure the precipitated fraction. After suspending the precipitated raction in 3.1 ml of water, 0.19 ml of 1 M Tris buffer (pH 9.2), centrifugation to recover the precipitated fraction and remove nost parts of contaminated proteins, cell components and the ike. The resulting precipitate was suspended in 4 ml of water and subjected to centrifugation to recover the precipitated raction containing h6T(1-163). The thus recovered precipitated fraction was suspended in 3.8 ml of water, and disintegrator and then subjected to centrifugation to recover A 0.6 g partion of frozen cells of the h6T(1-163)temperature for 40 minutes.

Just before the administration and on the next day of and 3

each of 6 animals of the control group in the same manner.

days after the completion of the administration, blood was collected from the eyeground to measure platelet counts using a microcell counter (F800, manufactured by Toa Iyo Denshi). In

days with a dose of 170 µl (having a total activity 22,000 measured by the rat CFU-MK assay system) per animal. As a control, 170 µl of DPBS was subcutaneously administered to

age), each group including 4 animals, daily for 5 consecutive

The TPO active fraction prepared above was

strong TPO activity (relative activity, 58,000) was found.

When this fraction was thoroughly dialyzed against IMDM culture medium and evaluated by the rat CFU-MK assay system,

final volume of a fraction (protein concentration, 2.18 mg/ml)

subjected to an in vivo assay. That is, the active fraction was

subcutaneously administered to ICR male mice (7 weeks of

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raction was analyzed by SDS-PAGE in the presence of a stirring, followed by 20 minutes of stirring at room solution was subjected to centrifugation to recover the supernatant fraction containing h6T(1-163), and a 5 ml portion 20 mM Tris buffer (pH 7.7) and then with 2.6 g of 20 to 50 mesh Dowex 1-x4 chloride form ion exchange resin, followed by 90 minutes of stirring. The ion exchange resin was removed using a glass filter to recover the resin-nonadsorbed fraction which was subsequently subjected to centrifugation to recover he supernatant fluid. The thus obtained supernatant fluid was applied to an SP Sepharose Fast Flow cation exchange column which has been equilibrated in advance with 20 mM Tris buffer (pH 7.7) and eluted with the same buffer by a linear gradient of rom 0 to 500 mM NaCl. When each of the eluted fractions of the SP Sepharose Fast Flow cation exchange column chromatography was thoroughly dialyzed against IMDM culture medium and evaluated by the rat CFU-MK assay system, strong rPO activity (relative activity, 19,000,000) was found in a raction eluted with a NaCl concentration of about 100 mM. This fraction was concentrated 1.6 fold (2.5 ml; protein concentration, 25 µg/ml) using an ultrafiltration unit (Ultra Free CL, nominal molecular cutoff of 5,000, manufactured by Millipore, catalog No. UFC4LCC25). When the thus concentrated educing agent, a band corresponding to TPO was detected as temperature to solubilize the contents. The resulting solution was mixed with 5 µl of 1% copper sulfate and stirred overnight (about 20 hours) at room temperature. The resulting of the supernatant was mixed with 5 ml of water and 10 ml of 0.2 ml of 1 M Tris buffer (pH 8) and 1 ml of 10% sodium Nlauroyl sarcosinate were added to the suspension with he main band (purity, 70 to 80%).

The TPO active fraction prepared by the above procedure was examined in an *in vivo* assay. That is, the active fraction was subcutaneously administered to ICR male

next day of the completion of the administration, blood samples were collected from the eyeground to measure containing 100 mM NaCl was subcutaneously administered in the same manner. Just before the administration and on the completion of the administration, in comparison with the than in the control group with a significant difference (p <0.01, Student t-test) between them. These results platelet counts using a microcell counter (F800, manufactured by Toa Iyo Denshi). In the TPO-administered group, platelet counts increased by a factor of about 1.73 in average after number prior to the administration and were 1.59 times higher demonstrated that the human TPO produced by E. coli and mice (8 weeks of age), each group including 4 animals, daily for 5 consecutive days at a dose of 100 µl (having a total activity of 47,500 measured by the rat CFU-MK assay system) oer animal. As a control, 100 µl of 20 mM Tris buffer (pH 7.7) prepared by the above procedure has the ability to increase platelet counts in vivo.

<Example 55>

Large scale culture of CHO cells

The large scale cultivation of the human TPO producing CHO cell line (CHO 28-30 cell, resistant to 25 nM MTX) which had been obtained by transfecting the human TPO expression plasmid pDEF202-hTPO-P1 into CHO cells in Example 32, was carried out as follows. The CHO cell line was cultured and proliferated in DMEM/F-12 culture medium (GIBCO) containing 25 nM MTX and 10% FCS. After the cells were detached with trypsin solution, 1 X 10⁷ cells were innoculated in a roller bottle (ex Falcon, Falcon 3000) containing 200 ml of the same medium therein and then cultured at 37·c at a rolling speed of 1 rpm for 3 days. After 3 days culture, the culture supernatant was removed by suction and the adherent CHO cells were then rinsed with 100

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containing neither 25 nM MTX nor 10% FCS was added to the bottle and the cells were cultured at 37°c at a rolling speed of 1 rpm for 7 days. After 7 days culture, the culture supernatant was recovered as a starting material for subsequent The above mentioned procedures were carried out in 500 roller bottles to obtain 100 L of the culture ml of PBS. 200 ml of the DMEM/F-12 medium (GIBCO) purification procedure. supernatant.

<Example 56>

Purification of human TPO from human TPO producing CHO cell

About 100 L of the serum free culture supernatant obtained in Example 55 was filtered through a concentrated on an ultrafiltration unit (PLTK Pellicon cassette, nominal molecular weight 30,000 cutoff, ex After the replacement of the solvent of the and Pefabloc SC (Merck), which are proteinase inhibitors, were added to the resulting aqueous solution at final concentrations of 1 mM and 0.35 mM respectively to obtain 3628 ml of the fraction of molecular weight 30,000 or greater (protein concentration: 1.60 mg/ml; total protein: 5805 mg; relative fraction was subsequently subjected to the Western blot analysis as described in Example 45. As a result, the presence between 66,000 and 100,000. In examining in more detail, it concentrate with pure water, p-APMSF (Wako Pure Chemicals) of TPO protein was revealed at a molecular weight ranging 0.22 µm filter to obtain its filtrate which was then was found that lower molecular weight TPO species than that of the above TPO were contained in the flow-through fraction. activity; 1,230,000; total activity: 7,149,000,000). Millipore).

The ultrafiltrate that contained the TPO with a molecular weight of not more than 30,000 was separately concentrated using an ultrafiltration unit (PLGC Pellicon

reveals that the low molecular weight TPO species were to obtain 1901 ml of the low molecular weight TPO fraction cassette, nominal molecular weight 10,000 cutoff, Millipore), (protein concentration: 0.36 mg/ml; total protein: 684 mg; relative activity: 245,500; total activity: 167,900,000). generated during cell culture.

Next, to 3614 ml of the fraction containing the TPO having a molecular weight of 30,000 or greater were insoluble substances were removed from the solution by added 764 g of ammonium sulfate and 144.5 ml of 0.5 M sodium citrate buffer (pH 5.5) to make 4089 ml of the solution containing 1.41 M (final concentration) ammonium sulfate and 17.7 mM (final concentration) sodium citrate buffer. After centrifugation, the clear solution was obtained.

Methyl HIC column (Bio-Rad, Catalog No. 156-0080; diameter 5 The clear solution was then applied to Macro-Prep cm, bed height 24.5 cm) which had been previously equilibrated ammonium sulfate at a flow rate of 25 ml/min. After the completion thereof, the elution was carried out with 20 mM sulfate. The eluted fraction was subsequently concentrated on with 20 mM sodium citrate buffer (pH 5.5) containing 1.2 M sodium citrate buffer (pH 5.5) containing 1.2 M ammonium an ultrafiltration unit (ex Filtron, Omega Ultrasette, nominal molecular weight 8,000 cutoff) to obtain a flow-through fraction F1 (4455 ml; protein concentration: 0.400 mg/ml; otal protein: 1780 mg; relative activity: 1,831,000).

ml; protein concentration: 0.969 mg/ml; total protein: 1411 mg; relative Next, 20 mM Na citrate (pH 6.0) as an eluting buffer was passed through the column to obtain an eluate which was then concentrated using the same ultrafiltration activity: 1,715,000). In the F2 the presence of a protein having unit (i.e., Omega Ultrasette, nominal molecular weight 8,000 a molecular weight from 66,000 to 100,000 was confirmed on cutoff) to collect a fraction F2 (1457

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SDS-PAGE. As a result of Western analysis, it was revealed hat the protein was TPO.

nominal molecular weight 8,000 cutoff; and Amicon, YM 3 elative activity: 5,558,000) which was then concentrated to 20 mM sodium citrate buffer (pH 6.0) at a flow rate of 15 out with 20 mM sodium citrate buffer (pH 6.0) containing 50 679 mg; relative activity: 88,830). Thereafter, the eluting buffer was replaced with 20 mM sodium citrate buffer (pH 5.4) protein concentration: 0.763 mg/ml; total protein: 710 mg; HIC. column (1443 ml) was applied to SP Sepharose Fast Flow bed height 12 cm) which has been previously equilibrated with ml/min. After the completion thereof, the elution was carried mM NaCl. The eluate was collected and named fraction F1 (3,007 ml; protein concentration: 0.226 mg/ml; total protein containing 750 mM NaCl to collect fraction F2 eluted (931 ml; 202 ml using ultrafiltration units (Filtron, Omega Ultrasette, Next, the F2 eluted from the Macro-Prep Methyl Pharmacia Biotech, Catalog No. 17-0729-01; diameter 5 cm, nembrane).

1,155,000), respectively. Thus, the fractionated TPO activity diameter 7.5 cm, bed height 100 cm) at a flow rate of 3 to obtain fractions F1 (585 ml; protein concentration: 1.00 mg/ml, total protein: 589 mg; relative activity: 4,118,000), F2 59.2 mg; relative activity 2,509,000), F3 (270 ml; protein concentration: 0.119 mg/ml, total protein; 32.1 mg; relative activity: 2,535,000) and F4 (720 ml; protein concentration: 3.0467 mg/ml; total protein: 33.6 mg; relative activity: (225 ml; protein concentration: 0.263 mg/ml; total protein: Next, the concentrated TPO activity-containing raction F2 (197 ml) was applied to a Sephacryl S-200HR gel iltration column (Pharmacia Biotech, Catalog No. 17-0584-05; 2,010-2,280 ml and 2,280-3,000 ml were separately collected Elution volumes 1,200-1,785 ml, 1,785-2,010 ml, nad a broad molecular weight range as determined nl/min.

Mestern blot analysis, it was found that F1 had a TPO molecule species with a molecular weight of 66,000 to 1,000,000 weight of 32,000 to 60,000; and F3 a TPO species with a molecular weight of 32,000 to 42,000. All the TPO molecules After SDS-PAGE of the fractions followed by predominantly; F2 a TPO molecule species with a molecular possessed TPO activity. Iltration.

On the basis of the result of N-terminal amino acid sequencing of the TPO molecule having a molecular weight of 66,000 to 100,000, we confirmed that the TPO molecule had the amino acid sequence of a human TPO gene encoded protein.

In addition, the TPO molecule having a molecular glycanase (ex Genzyme, Catalog No. 1472-00), neuraminidase SDS-PAGE analysis. As a result, the polypeptide portion of the TPO was found to have a molecular weight of about 36,000 as expected from its theoretical molecular weight and to be a weight 66,000-100,000 was subjected to enzymatic digestion experiments using the glycosidase enzymes consisting of N-Nakarai-Tesqu, Catalog No. 242-29SP), endo-a-N-100453) and O-glycosidase (Boehringer Mannheim Biochemica, Catalog No. 1347101) singly or in combination, and then to acetylgalactosaminidase (Seikagaku Kogyo, Catalog No. glycoprotein with both N- and O-linked sugar chains.

cExample 57>

Purification of human TPO from human TPO producing CHO cell line

was added 211.4 g of ammonium sulfate, after which the mixture was filtered through a 0.2 µm filter (ex Gelman he CHO cells obtained in the same manner as in Example 55 Science, Catalog No. 12992). The obtained filtrate was applied to Macro-Prep Methyl HIC column (Bio-Rad, Catalog No. 156-1081, diameter 50 mm, bed height 90 mm) which had been (1) To 1 L of the serum free culture supernatant of

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then applied to a reverse phase Vydac C4 column (The 5.6) containing 1.2 M ammonium sulfate at a flow rate of 15 ml/min. After the completion thereof, the elution was carried containing 1.2 M ammonium sulfate. The eluted fraction was bed height 250 mm) at a flow rate of 0.75 ml/min. After the min, the elution was carried out with a 66-min linear gradient of from the development solvent A to 10 mM Tris buffer (pH 6.4) containing 94% ethanol (referred to as "development As a result of SDS-PAGE analysis, the molecule having a molecular weight 65,000-100,000 which was expected to be TPO and which corresponded to the fraction of retention time 68-72 min after addition of the sample, appeared as a single previously equilibrated with 20 mM sodium acetate buffer (pH out with 450 ml of 20 mM sodium acetate buffer (pH 5.6) Separations Group, Catalog No. 214BTP54, diameter 4.6 mm, column was washed with 10 mM Tris buffer (pH 6.4) containing 5% ethanol (referred to as "development solvent A") for 15 solvent B"). The chromatogram obtained is shown in Fig. 15. band on the SDS-PAGE gel (see Fig. 16). Western analysis urther revealed that the obtained protein is TPO.

The N-terminal amino acid analysis of the protein sample revealed that the protein had the amino acid sequence of a human TPO gene encoded protein.

Example 58>

human TPO-expression within insect cells Preparation of recombinant virus for

Sene DNA purification kit. The purified DNA was then ligated digested with restriction enzymes EcoRI and Notl and then subjected to 1% agarose gel electrophoresis to obtain an about 1200-bp band which was subsequently purified using Prep-Ato a transfer vector pVL 1393 (Invitrogen) pretreated with the same restriction enzymes, followed by transformation into The plasmid pHTP1 prepared in Example 30 was

Competent-High E. coli DH5 (Toyo Boseki). From the colonies obtained was selected a clone pVL1393/hTPO which contained a full length coding region of human TPO cDNA. The plasmid DNA was then prepared from the clone by the method described Laboratory Press, 1989) followed by transfection into insect cell Sf 21 (Invitrogen) using BaculoGoldTM Transfection kit (Farmingen), and the transfected cells were cultured in Sf-900 medium (Lifetechnology) at 27.c for 4 days to recover the supernatant containing the virus. The supernatant was then diluted to 1:109 \sim 1:105, and about 7 X 105 Sf 21 cells were infected with 1 ml of the diluted supernatant at 27 c for 1 hr in a 35-mm (in diameter) shale. After removal of the supernatant, 1% hot agarose in Sf-900 medium was placed into the culture and allowed to solidify, followed by 6-day cultivation at 27 c in a moisturized atmosphere. A single plaque formed was picked up, from which the virus was released in 200 µl of Sf-900 medium. Sf 21 cells were infected with the obtained viral clone in a 24-well plate to proliferate the virus. A portion of the virus-containing liquid rom the single plaque was treated with phenol/chloroform followed by ethanol precipitation to recover the viral DNA which was then used as a template to carry out PCR using primers specific for human TPO cDNA. The recombinant virus containing human TPO cDNA was selected based on amplification of the specific DNA fragment. Then, Sf 21 cells vere infected with the supernatant containing the recombinant in Molecular Cloning (Sambrook et al., Cold Spring Harbor irus carrying human TPO cDNA to proliferate the virus.

<Example 59>

Expression of human TPO in Sf 21 insect cells and identification of TPO activity

Sf 21 cells were cultured in a 175 cm² culture lask to about 80% confluence was attained, followed by

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infection with the human TPO cDNA carrying recombinant virus prepared in Example 58 at 27 c for 1 hr, after which the obtained cells were cultured in Sf-900 medium at 27 c for 4 days to recover the supernatant of the culture. The obtained culture supernatant was substituted by IMDM culture medium on NAPTM-5 column (Pharmacia). Significant TPO activity in the resultant supernatants was detected in a dose-dependent manner in both rat CFU-MK assay and M-07e assay. Also, the recombinant human TPO expressed in Sf 21 cells was identified by means of the Western analysis as described in Example 45.

<Example 60>

clone oCFM536/h6T(1-163) carrying a human TPO base auroyl sarcosinate and cupric sulfate, and purification of the sequence via its expression in E. coli. by using sodium N-Refolding of variant human TPO, h6T(1-163) derived from h6T(1-163) Thirty g of the frozen h6T(1-163)-producing ecombinant microorganism prepared in Example 43 was suspended in 300 ml of water, disrupted in a high pressure disrupting apparatus (10,000 psi, Rannie High Pressure Laboratory), and then centrifuged to recover a sedimented fraction. The sediment fraction was suspended in 90 ml of water to which water was then added to the total amount of 50 ml with stirring. Subsequently, to the mixture was added 9 ml of 1 M Tris buffer (pH 9.2), 540 µl of 1 M DTT, 1.8 ml of 0.5 M EDTA and 18 ml of 10% sodium deoxycholic acid with stirring, followed by 30 min-stirring at room temperature. The sedimented fraction was recovered by centrifugation and most of the contaminated proteins and microorganism components were removed. To the sediment was added 180 ml of 5 mM DDT to obtain a suspension which was then centrifuged to recover the sediment fraction containing h6T(1-

To the resultant sediment was added 300 ml of water to a suspension to which water was then added to the iquid amount of 570 ml with stirring. Thirty ml of 1 M Tris ouffer (pH 8), 150 ml of 10% sodium N-lauroyl sarcosine were added to the mixture at room temperature with 20-min stirring in order to solubilize the h6T(1-163). To this was urther added 750 µl of 1% cupric sulfate and the mixture was stirred overnight (about 20 hr) at room temperature. After the Tris buffer (pH 9.2) containing 0.1% polysorbate 80 to recover fraction was adjusted to 7.2 with hydrochloric acid, applied to SP Sepharose Fast Flow cation exchange column (ID 5 cm X 10 0.1% polysorbate 80, and then eluted with the linear gradient fractions were subjected to SDS-PAGE analysis to collect a TPO fraction to which trifluoroacetic acid was then added to a to Capcell Pak 5 µm 300 A C1 column (ID 2.1 cm X 5 cm X 2, centrifugation, to 750 ml of the supernatant was added 750 ml by addition of 3 ml of polysorbate 80 (Nikko Chemicals) with stirring. To the obtained solution was added 600 g of Dowex by 90-min stirring at room temperature. Following recovery of the non-adsorbed fraction using a glass filter, the ionexchange resin was washed with 750 ml of 20 mM Tris buffer fraction was adjusted to pH 9.2 with 2 N sodium hydroxide after which it was applied to Q Sepharose Fast Flow anionexchange column (ID 5 cm X 10 cm) equilibrated with 20 mM a non-adsorbed fraction. The pH of the obtained non-adsorbed cm) equilibrated with 20 mM Tris buffer (pH 7.2) containing from 0 M to 500 mM NaCl in the same buffer. The eluted concentration of 0.1%. After the solution obtained was applied Shiseido), a reverse phase HPLC was carried out by the linear gradient elution method with increasing 1-propanol supernatant fraction containing h6T(1-163) was recovered by of water and 1,500 ml of 20 mM Tris buffer (ph 7.7) followed 1-X4 (20-50 mesh, chloride form) ion-exchange resin followed (pH 7.7). The combined solution of the non-adsorbed and wash

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analyzed on SDS-PAGE in the absence of a reducing agent. As a HPLC column in the similar manner. The resultant three TPO (see Fig. 17), were analyzed on SDS-PAGE in the absence of a concentration in 0.1% trifluoroacetic acid. The eluate was result, three fractions were fractionated based upon the elution position in the reverse phase HPLC, each fraction of Each fraction diluted was applied to the above reverse phase ractions which were designated as Fr. Sta, Fr. Stb and Fr. Stc depending in the order of elution on the reverse phase HPLC reducing agent. As a result, a single band was detected at a which was diluted three times with 0.1% trifluoro-acetic acid. molecular weight of about 18 kDa (Fr. S-a), about 19 kDa (Fr. S-b) or 18 kDa (Fr. S-c) (see Fig. 18). Following their amino acid analyses, each of the amino acid compositions determined was almost consistent with the corresponding theoretical he results of N-terminal amino acid analysis showed that The protein amount of each raction determined by amino acid analysis was 0.64 mg (Fr. Sa), 1.81 mg (Fr. S-b) or 3.49 mg (Fr. S-c). After full dialysis of each fraction against IMDM medium, M-07e assay was carried out. As a result, the TPO relative activity of each fraction was about 1,620,000 (Fr. s-a), 23,500,000 (Fr. S-b) or value estimated from the sequence information. Additionally, hey were the expected sequences. 746,000,000 (Fr. S-c).

<Example 61>

and cystein-cystin, and purification of the sequence via its expression in E. coli. by using quanidine Refolding of variant human TPO, h6T(1-163) derived rom clone pCFM536/h6T carryinga human TPO base (ET(1-163) Fifty g of the h6T(1-163)-producing frozen recombinant microorganism prepared in Example 43 was added to 500 ml of water and suspended, disrupted using a high

Rannie High sediment fraction. The sediment fraction was suspended in M Tris buffer (pH 9.2), 900 µl of 1 M DTT, 3 ml of 0.5 M EDTA and 30 ml of 10% sodium deoxycholic acid were added and suspended, after which the sedimented fraction containing n6T(1-163) was recovered by means of centrifugation. The recovered sedimented traction containing h6T(1-163) was suspended in water in a total volume of 104 ml. To the suspension was added 20 ml of 1M Tris buffer (pH 8.5) and then 376 ml of 8 M guanidine hydrochloride with stirring, and to solubilize the h6T (1-163). To this was added 2,500 ml of then 2,000 ml of 20 mM Tris buffer (pH 8.5) containing 1 M. 5 mM cystein and 0.5 mM cystin. The thus obtained solution was incubated overnight at 4 c and then centrifuged to recover 16T(1-163) in the supernatant which was then concentrated using Prep Scale UF cartridge PLDC ultrafiltration membrane (Millipore). The buffer of the concentrate was replaced by 20 was applied to Q Sepharose Fast Flow anionic exchange column containing 0.1% polysorbate 80, and then eluted with a linear gradient from 0 M to 500 mM NaCl in the same buffer, by which he fraction (240 ml) eluted at about 20 mM - about 150 mM water and its liquid amount was then adjusted to 250 ml by stirred at room temperature for 30 min. After centrifugation, he sediment fraction was recovered while removing most of he contaminated proteins, microorganism components and the ike. To the sediment was added 300 ml of 5 mM DTT and subsequently stirred at room temperature for 10 min in order 20 mM Tris buffer (pH 8.5) containing 0.1% polysorbate 80 and guanidine hydrochloride with stirring, followed by addition of mM Tris buffer (ph 9.2) containing 0.1% polysorbate 80 until reaching a final volume of 1,000 ml. The obtained solution ID 5 cm X 10 cm) equilibrated with 20 mM Tris buffer (pH 9.2) addition of water with stirring. To this, thereafter, 15 ml of 1 to recover pressure disrupting apparatus (10,000 psi, Pressure Laboratory), and then centrifuged

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VaCI was recovered. The resultant fraction was diluted four

gradient from 0 M to 500 mM NaCl in the same buffer. The imes with 20 mM Tris buffer (pH 7.2) to attain a total volume applied to SP Sepharose Fast Flow cation exchange column (ID containing 0.1% polysorbate 80, and then eluted with a linear of 960 ml, and then adjusted to pH 7.2 with acetic acid, 5 cm X 10 cm) equilibrated with 20 mM Tris buffer (pH 7.2)

sluate was analyzed on SDS-PAGE to collect a TPO fraction. To

the TPO fraction was added trifluoroacetic acid until its final Pak 5 µm 300A C1 column (ID 2.1 cm X 5 cm X 2, Shiseido), and volume of 0.1%. The resultant solution was applied to Capcell

everse phase HPLC was subsequently conducted by linear gradient elution method with increasing 1-propanol concentration in 0.1% trifluoroacetic acid. The eluate was analyzed by SDS-PAGE in the absence of a reducing agent and

position in the reverse phase HPLC. The two TPO fractions ractionated into two fractions depending on the eluted were designated as Fr. G-a and Fr. G-d based on their order

eluted in the reverse phase HPLC (see Fig. 17). Each fraction agent. As a result, a single band was detected at a molecular weight of about 18 kDa (Fr. G-a) or about 32 kDa (Fr. G-d) (see was then analyzed on SDS-PAGE in the absence of a reducing

Fig. 18). Furthermore, the SDS-PAGE analysis of the above ractions in the presence of a reducing agent showed that in both the fractions a single band was detected at a molecular weight of about 20 kDa. The result of the amino acid analysis

compositions thereof was almost consistent with the corresponding theoretical value estimated from the sequence The protein amount of each fraction determined from the of the fractions showed that each of the amino acid nformation. Additionally, the results of the N-terminal amino acid analysis showed that they were the expected sequences.

results of the amino acid analysis was 2.56 mg (Fr. G-a) or

1.16 mg (Fr. G-d). Following full dialysis of each fraction

about against IMDM medium, M-07e assay was carried out. As result, the fraction had TPO relative activity of 3,960,000 (Fr. G-a) or of about 7,760,000 (Fr. G-d).

<Example 62>

expression of a partial length human TPO (amino acids 1 to 163) (hereinafter referred to as "hTPO163") within CHO cells Construction of recombinant vector pDEF202-hTPO163 for

gene (Amp^r), in which the hTPO163 cDNA is linked at a site electrophoresis. The fragment was ligated by T4DNA ligase (Takara-Shuzo) with an hTPO163 cDNA which had been prepared by treating a plasmid pEF18S-hTPO163 containing the hTPO163_cDNA encoding the amino acids -21 to 163 (SEQ ID an origin of replication of SV40, a human elongation factor 1α-promoter, an SV40 early polyadenylation site, a murine DHFR minigene, an origin of replication of pUC18 and a \(\beta^{-1}\)actamase The vector pDEF202 constructed in Example 31 was ecovery of a larger vector fragment using agarose gel NO:13) with restriction enzymes EcoRI and Spel, to give an expression vector pDEF202-hTPO163. This plasmid contained reated with restriction enzymes EcoRI and Spel followed by downstream of the human elongation factor 1- α -promoter.

Example 63>

Expression of hTPO163 in CHO cells

A CHO cell line (dhfr-strain, Urlaub and Chasin, Proc. Natl. Acad. Sci. USA, ZZ, p 4216, 1980) was grown in α minimum essential medium ($\alpha ext{-MEM}(ext{-})$ with thymidine and (Falcon), and then transformed with a pDEF202-hTPO163 hypoxanthine) containing 10% bovine fetal serum in 6 cm dish plasmid by the transfectum method (Seikagaku Kogyo K.K.).

prepared in Example 62 was mixed with 240 µl of 0.3 M NaCl Briefly, 10 µg of the plasmid pDEF202-hTPO163 a mixture of 20 µl of transfectum and 220 µl of H2O. then with

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exchanged with the fresh selection medium at intervals of two his DNA solution was added dropwise to the dish followed by culture for 6 hr in a ${\sf CO}_2$ incubator. The medium was removed rom the dish which was then washed twice with $\alpha\text{-MEM}(\text{-})$ ollowed by addition of 10% DMSO containing $\alpha\text{-MEM}(\text{--})$ before ncubation for 2 minutes at room temperature. Thereafter, to he dish was added 10% dialyzed bovine fetal serum-containing non-selection medium (a-MEM(-) with hypoxanthine and hymidine) followed by two day culture, then selection in 10% Jialized bovine fetal serum-containing selection medium (α -MEM(-) without hypoxanthine and thymidine). The selection was conducted by trypsinizing the cells, dividing them into ive 10 cm dishes or twenty 24 well dishes per the above 6 cm dish, and continuing to culture while the medium was days. The supernatants from plates or wells were assayed for IPO activity using Ba/F3 assay, thereby the TPO activity being observed. The cells in which TPO activity was observed in the culture supernatants were then transferred to fresh plates or wells after divided to a cell concentration of 1:15 with the subsequently recultured for growth and cloning of cells selection medium containing 25 nM methotrexate, esistant to methotrexate.

Alternatively, the transformation of the CHO cells can be carried out by co-transfection of pEF18S-hTPO163 and pMG1 into the CHO cells.

applicant on January 31, 1995, under the accession No. FERM BP-4989, at the National Institute of Bioscience and Human CHO strain (CHO-DUKXB11) transfected with the plasmid pDEF202-hTPO163 has been deposited by the present Fechnology, Agencey of Industrial Science and Technology Ministry of International Trade and Industry, Japan.

arge-scale culture of CHO cells Example 64>

CHO cell line (CHO 109 cells, obtained by the above-mentioned in 10% dialyzed bovine fetal serum-containing medium [α -MEM" without hypoxanthine and thymidine]) which had been obtained by transfection of the TPO163-expression plasmid pDEF202-hTPO163 into CHO cells in Example 63 was carried out as follows. The CHO 109 cells After the cells were harvested (or trypsinized) using a trypsin solution, ten millions cells were inoculated into a Falcon roller bottle (Falcon 3000) containing 200 ml of the same medium and then cultured at a rolling speed of 1 rpm at 37.c for 3 days. The culture medium was removed under suction and the surface of the cell culture was rinsed with 100 ml of PBS. Then, to the cell culture was added 200 ml of DMEM/F-12 medium (GIBCO) without 10% FCS followed by 7 day culture at 37.c at 1 rpm. The culture supernatant harvested was used as a starting material for the subsequent purification step. The similar procedures were conducted in 300 roller bottles to The large-scale culture of the hTPO163-producing were grown in DMEM/F-12 medium (GIBCO) with 10% FCS. rield 60 L of the serum-free culture supernatant. selection

<Example 65>

Purification of hTPO163 from hTPO163-producing CHO cell line

weight 10,000 cutoff) to obtain a concentrated fraction (600 ml, 11.2 mg protein/ml, total protein 6430 mg). By Western the presence of hTPO163 protein expressed at the range of an (1) 60 L of the serum-free culture supernatant obtained in Example 64 was filtered through a 0.22 µm iltration filter to collect a filtrate which was then concentrated using an ultrafiltration unit (Filtron, molecular analysis of this fraction using the anti-HT1 peptide antibody, apparent molecular weight from 20,000 to 26,000 was shown.



(Pharmacia-Biotech, Catalogue No. 17-0032-02; diameter 10 phosphate buffer (pH 6.8) to obtain a protein fraction F1 (938 ml, protein concentration 4.9 mg/ml, total protein 4594 mg) as cm and bed height 30 cm) pre-equilibrated with 10 mM sodium The resulting concentrated culture supernatant (537 ml) was treated on a Sephadex G-25 Fine column a solution in 10 mM sodium phosphate buffer (pH 6.8).

The column was applied at a flow rate of 15 ml/min, to a SP 17-0729-01; diameter 5 cm and bed height 12 cm) preequilibrated with 10 mM sodium phosphate buffer (pH 6.8) ollowed by elution with 10 mM sodium phosphate buffer (pH 5.8) then the same buffer containing 10% ethanol. The eluates were combined as a fraction F1 (1608 ml, protein concentration 2.13 mg/ml, total protein 3426 mg). Then, the second elution was carried out with 10 mM sodium phosphate buffer (pH. 6.8) containing 750 mM NaCl and 25% ethanol to collect a main hTPO163 eluate of fraction F2 (651 ml, protein This protein fraction (929 ml) from Sephadex G-25 Sepharose Fast Flow column (Farmacia-Biotech, Catalogue No. concentration 1.67 mg/ml, total protein 1087 mg).

10 mM sodium acetate buffer, pH 6.7) plus 50% of solvent B 55% solvent A plus 45% solvent B until unadsorbed substances were almost completely eluted, after which the elution at a tow rate of 1.5 ml/min was carried out according to the ethanol solution. Insoluble materials which resulted from the addition of ethanol were removed by centrifugation. The supernatant was injected into a SOURCE 15RPC column Pharmacia-Biotech, Catalogue No. 17-0727-02; diameter 2 cm and bed height 20 cm) pre-equilibrated with 50% of solvent A (10 mM sodium acetate buffer, pH 6.7, containing 90% ethanol) at a flow rate of 2 ml/min. Then the column was washed with To the TPO activity-containing fraction F2 (200 ml) from SP Sepharose Fast Flow column were added ethanol and purified water to prepare 300 ml of 45% (final conc.)

35 minutes. The fractions were collected at intervals of 5 ourified hTPO163 with an apparent molecular weight from oroteins, the hTPO163 having a higher molecular weight was collowing elution profile: 50% B for 5 minutes; linear gradient from 50% B to 100% B over 140 minutes; and then 100% B for minutes (corresponding to 7.5 ml volume). All the fractions examining an elution range of hTPO163. As a result, the highly approximately 20,000 to 26,000 was found to be eluted at the range from 66% to 87% ethanol. Among these hTPO163 eluted earlier from the reverse phase column, suggesting that were subjected to SDS-PAGE then to Western analysis for the glycosylated hTPO163 molecule has an increased nydrophilicity.

application, the elution fractions were collected in an amount of 6 ml each (i.e. every 4 min). As a result, the hTPO163 was hTPO163 fraction (90 ml) eluted at the range from 68% to This concentrate was subsequently applied to a Superdex 75 pg column (Pharmacia-Biotech, Catalogue No. 17-1070-01; mM sodium phosphate buffer (pH 6.8) containing 10% ethanol at a flow rate of 1.5 ml/min. From the time of 60 minutes after eluted in the fractions from tube number 16 to at least tube number 31 as determined by SDS-PAGE. This elution position is corresponding to the range of a molecular weight from approximately 44,000 to approximately 6,000, as determined by gel filtration using a standard molecular weight marker Additionally, these hTPO163 molecules appeared to be eluted in the order of decreasing degree of glycosylation. All the To 88.8 ml of the hTPO163 elution fraction (i.e. 86.5% ethanol) from SOURCE 15RPC column was added CHAPS, and the mixture was concentrated and washed to obtain 2.5 ml of a concentrate containing about 5% ethanol and 4 mM CHAPS. diameter 2.6 cm and bed height 60 cm) equilibrated with 10 mixture of Bio-Rad, Gel Filtration Standard, Catalogue No. 151-1901; and Calbiochem, insulin, Catalogue No. 407696)

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hTPO163 species of the range from tube numbers 16 to 31 (elution volume: 180 to 276 ml) were collected as fraction FA. In addition to the fraction FA, the fractions of the tube Nos. 16 to 18 (elution volume: 180 to 198 ml), 19 to 24 (elution volume: 198 to 234 ml) and 25 to 31 (elution volume: 234 to 276 ml) were separately collected and designated as fractions FH, FM and FL, respectively. Also, the fraction FA was prepared by combination of the portions of the fractions FH, FM and FL. The above is shown in Fig. 19.

the N-terminal Ser residues could not be identified. This illustrated in more detail. The N-terminal amino acid sequences of the hTPO163 species obtained above were determined by the same method as in Example 1, but most of suggests that an O-linked sugar is added to the N-terminal Ser. Furthermore, the sequence following the N-terminal Ser was confirmed to be an amino acid sequence expected from the gene sequence of hTPO. The concentrations of the hTPO163 proteins in the FH, FM, FL and FA were 10.2 ng/ml, 6.2 ng/ml, 0.84 ng/ml and 3.2 ng/ml as determined by amino acid analysis (AccQ. Tag method, Wasters), respectively, provided that the sugar chain. These fractions (100 ng each) were subjected to SDS-PAGE under unreducing conditions using a multigel 15/25 (Dai-ichi Kagaku Yakuhin, 15 to 25% precast polyacrylamide these fractions was found to contain highly pure hTPO163. The and FA, as calculated using DPCIII molecular weight marker (Dailchi Pure Chemicals) as a standard under the reducing and FA from the Superdex 75 pg column set forth in (1) will be concentrations were of the peptide moieties not containing any gel) or under reducing conditions using DTT, followed by silver-staining (Dailchi Pure Chemicals). As a result, each of apparent molecular weights of the hTPO163 in the FH, FM, FL conditions, were 24,000 to 21,500, 23,000 to 21,000, 23,000 (2) Next, the hTPO163 elution fractions FH, FM, FL 20,500 and 23,500 to 20,500, respectively (see Fig. 20).

Additionally, the apparent molecular weights of the hTPO in Broad Range: Catalogue No. 161-0319) under the reducing espectively. The heterogeneity among the molecular weights of FH, FM, FL and FA may be due to the heterogeneity of the Oand then analyzed on SDS-PAGE. As a result, in all the evealing that the heterogeneity of the molecular weight of hTPO is mainly due to the heterogeneity of the amount of sialic PAGE standard (Bio-Rad, Biotynylated SDS-PAGE Standards, conditions on Western analysis, were 26,000 to 22,000, 25,500 to 22,000, 26,000 to 21,000 and 26,000 to 21,000, acid in the sugar chains coupled with the hTPO163 protein and that the hTPO163 obtained was expressed as a glycoprotein in he FH, FM, FL and FA, as calculated using biotin-labelled SDSinked sugar chains. Therefore, each fraction of the FH, FM, FL and FA was digested with neuraminidase (Neuraminidase, Nacalai tesque, Catalogue No. 242-29SP), reduced with DTT, fractions, the apparent molecular weight was around 19,000, CHO cells.

were assayed for their <u>in vitro</u> activity by M-07e assay system. As a result, the relative specific activity were 511,000,000, 775,000,000, 1,150,000,000 and 715,000,000/mg hTPO163 protein (weight of the peptide moiety not containing any sugar weight).

<Example 66>

Construction of E. coli vector for expression of human TPO (amino acids 1 to 332) in which Lys is added at the position-1 and Met. at the position-2 respectively (hereinafter referred to as "hMKT(1-332)"), and expression of the hMKT(1-332)

To express the full length amino acid sequence of human TPO in <u>E. coli</u>, the use codons of from amino acid 164 to amino acid 332 were changed to <u>E. coli</u> preference codons as described below.



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Table 5

21 : S-CTDCDCBAACDGTACCAGOGGCCTGCTGGAAACCAACTTTACCGCGAGS

(SEQ ID NO:130)

2:5-GGTAAAGTTGGTTTDCAGCAGGCOGCTGGTACGGGTTCGGGAGCTCGT-3

23:5-03030STACCACCGGCCAGCGGCCTGCTGAAATGGCCAGCAGGGCTTTCGT-3

(SEQ ID NO:132)

(SEQ ID NO:131)

24:5-AGCOCTGCTGCCATTTCAGCAGGCCGCTGCCGGTGGTACGCGCGCTCGC3

(SEQ ID NO:133)

25.5-GDGAAAATCCCCGGCCTGCTGAACCAGACCAGCCGTAGCCTGGATCAGAT:3

(SEQ ID NO:134)

28:5-ATDCAGGCTACGGCTGGTTCAGCAGGCCCGGGATTTTCGCACGAA-3

SEQ 1D NO:135)

SEQ ID NO:136) Z7:S-COCGGGGCTATICTGAACCGTATCCATGAACTGCTGAACGGCACCCGTG-3

28:5-GTGCCGTTCAGCAGTTCATGGATACGGTTCAGATAGCCCGGGATCTG-3

SEQ ID NO:137)

28:5-GOCTIGT TICOGGGGCCGAGCCGTTCGCACCCTGGGCCGCCGCCGATATCAG3

(SEQ ID NO:138)

SEQ ID NO:139) 30:5-ATEODGGCGCCCAGGGTTGCGACGGCTTCGGGCCCCCGGAAACAGGCCCACGGG

SEQ 1D NO:140)

31:SATCAGOTOTGGCACCAGOGATACOGGCAGOOTGOOGCOGAACOTGCAGOC3

SEQ ID NO:141) 22:5-CAGGITTCGGCCGCCAGGCTGCCGGTATCGCTGGTGCCCAGAGCTGATATCCG3

33:5-GGGCTATAGCCCGAGCCCATCCGCCCGACCGGCCAGTATACCCTGTT-3

34:5-GGTATACTIGGCOGGTCGGCGGATGGGTCGGGCCTCGGGGCTATAGCCCGGGCTG3

SEQ ID NO:142)

\$5:5-TOOGCTGOCGCCGACOCTGCCACCCGCGTCATTCAGCTGCATCCGCTGC3

(SEQ ID NO:144)

37:5-TECOCEGATECCEACECCECCACCOCGACCAGCCCAGCCCAGCCAG

38:5Accacoccecotestossestrossestroscoscotossatrosscasoascas

(SEQ ID NO:148)

(SEQ ID NO:147)

39:5-CCAGCTATACCCATAGCCAGAACCTGAGCCAGGAAGGCTAATGAAGCTTGA3

40:5-CTTCATTAGCCTTCCTGGCTCTGGCTATGGGTATAGCTGGTGTTC-3

(SEQ ID NO:149)

(SEQ ID NO:151) (SEQ ID NO:150) 41:5-ACGAGCTCCCGAACCGTACCA-3" 42:5-CTGATATCCGGCGCGCCAGG3

(SEQ ID NO:152) (SEQ ID NO:153) 44:5-TCAAGCTTCATTAGCCTTCCT-3" 43:5-CGGATATCAGCTCTGGCACCA-3

and 36; 37 and 38; or 39 and 40, were phosphorylated in a The synthetic oligonucleotides: 21 and 22; 23 and 24; 25 and 26; 27 and 28; 29 and 30; 31 and 32; 33 and 34; 35 acetate, 50 mM K-acetate using T4 kinase (Pharmacia) in a same tube. Then 1/10 volumes of a solution of 100 mM Tis/HCI (pH 7.5), 100 mM MgCl₂, 500 mM NaCl was added to he reaction mixture, boiled for 3 minutes in a water bath, and allowed to cool to form a double stranded DNA. Four sets of the double stranded DNAs, i.e. oligonucleotides 21 and 22/23 and 24 (combination-A); oligonucleotides 25 and 26/27 and 28 (combination-B); oligonucleotides 31 and 32/33 and 34 combination-C); and oligonucleotides 35 and 36/37 and 38/39 and 40 (combination-D), were separately ligated with a DNA igation kit (Takara-Shuzo), and thereafter the combination-A was ligated together with the combination-B and similarly the and ligation-2 respectively. Using the ligations-1 and -2 as 11 and -42 for ligation-1, as well as the oligonucleotides-43 and -44 for ligation-2, were employed as primers. The solution of 0.1 mM ATP, 10 mM Tris-acetate, 10 mM Mgtemplates, PCR were carried out wherein the oligonucleotidescombination-C with the combination-D to produce ligation-1 same tube.



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purification using a Prep-A-Gene DNA Purification Kit to ecover about 240-bp and 250-bp fragments respectively. The ollowed by electrophoresis on 2% agorose gel and then wo fragments thus obtained were subcloned into pBluescript II KS+ (Stratagene) predigested with SacI and HindIII (E. coli DH5 was employed as a host.). Of the resulting colones, the clone having the base sequence shown in Table 6 was selected products from the PCR of the ligations-1 and -2 were digested with Sacl and EcoRV and with EcoRV and HindIII respectively, through sequencing and named pBL(SH)(174-332) (see SEQ ID

Table 6

CIC COG AAC CGI ACC AGC GGC CIG GAA ACC AAC III ACC GCG AGC GCG AAA ATC CCG GGC CTG CTG AAC CAG ACC AGC CGT AGC CTG GAT CAG Ala Lys Ile Pro Gly Leu Leu Asn Gln Thr Ser Arg Ser Leu Asp Gln Leu Pro Asn Arg Thr Ser Gly Leu Leu Glu Thr Asn Phe Thr Ala Ser GCG CGT ACC ACC GGC AGC GGC CTG CTG AAA TGG CAG CAG GGC TTT CGT Ala Arg Thr Thr Gly Ser Gly Leu Leu Lys Trp Gln Gln Gly Phe Arg 220 210 174 130

ATC CCG GGC TAT CTG AAC CGT ATC CAT GAA CTG CTG AAC GGC ACC CGT ile Pro Gly Tyr Leu Asn Arg Ile His Glu Leu Leu Asn Gly Thr Arg GGC CTG III CCG GGC CCG AGC CGI CGC ACC CTG GGC GCG CCG GAT ATC 3ly Leu Phe Pro Gly Pro Ser Arg Arg Thr Leu Gly Ala Pro Asp Ile 240

AGC TOT GGC ACC AGC GAT ACC GGC AGC CTG CCG AAC CTG CAG CCG Gly Thr Ser Asp Thr Gly Ser Leu Pro Pro Asn Leu Gln Pro Ser Ser (

36C TAT AGC COG AGC COG ACC CAT CCG COG ACC GGC CAG TAT ACC CTG Gly Tyr Ser Pro Ser Pro Thr His Pro Pro Thr Gly Gln Tyr Thr Leu

III CCG CTG CCG CCG ACC CTG CCG ACC CCG GTG GTT CAG CTG CAI CCG Pro Leu Pro Pro Thr Leu Pro Thr Pro Val Val Gln Leu His Pro Phe

CTG CCG GAT CCG AGC GCG ACC CCG ACC AGC CCG CTG Leu Pro Asp Pro Ser Ala Pro Thr Pro Thr Ser Pro Leu 9 Len

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cts aac acc tat acc cat agc cag aac cts agc cag gaa ggc taa Leu Asn Thr Ser Tyr Thr His Ser Gln Asn Leu Ser Gln Glu Gly 330 320

TGA AGC TTG A

Next, the pBL(XH)h6T(1-163) prepared in Example 42 as a template was subjected to PCR in the presence of the following synthetic oligonucleotides 45 and 46 as primers, and electrophoresis on 6% polyacrylamide gel to recover an about the PCR product was digested with BamHI and Sacl followed by 160-bp fragment from the gel.

(SEQ ID NO:155) 45: 5-AAGGATCCGAACGCTATCTTCCTG-3" 46:5-GGGAGCTCGTTCAGGGTCAGAACCAGA GAGGTACGAGACGGAACAGCAGTGGTTGG-3(SEQ ID NO:156)

Also, the pBL(SH)(174-332) was digested with Sacl and HindIII and the digest was then purified using the Prep-A-Gene DNA Purification Kit to give an about 480-bp fragment. The two fragments prepared above were subcloned into pBluescript II KS+ (Stratagene) predigested with BamHI and HindIII (E. coli OH5 was employed as a host.).

Of the resulting clones, the clone having the base sequence shown in Table 7 was selected through sequencing and named pBL(BH)(123-332) (see SEQ ID NO:157).

Table 7

G GAT CCG AAC GCT ATC TTC CTG TCT TTC CAG CAC CTG CTG CGT GGC

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Asp Pro Asn Ala Ile Phe Leu Ser Phe Gln His Leu Leu Arg Gly 130 123

Lys Val Arg Phe Leu Met Leu Val Gly Gly Ser Thr Leu Cys Val Arg AAA GIT CGI IIC CIG AIG CIG GIT GGC GGI ICI ACC CIG IGC.GIT

GGG GCG CCA ACC ACT GCT GTT CCG TCT CGT ACC TCT CTG GTT CTG Arg Ala Pro Pro Thr Thr Ala Val Pro Ser Arg Thr Ser Leu Val Leu

160

Thr Leu Asn Glu Leu ACC CTG AAC GAG CTC

170

174

with the same restriction enzymes to produce a clone expression vector was employed as a transformant for the The expression plasmid pCFM536/hMKT(1-332) contains a DNA together to obtain a clone pBL(XH)h6T(1-334). Further, the was subsequently ligated with a pBL(XH)h6T(1-334) digested pBL(XH)hMKT(1-334). After the pBL(XH)hMKT(1-334) was which codes for the mutation type human TPO amino acids 1-332 was recovered and purified, and then cloned into pCFM536 pCFM536/hMKT(1-332), and the E. coli strain carrying this expression of the mutation type protein hMKT(1-332) protein. The pBL(XH)h6T(1-163) prepared in Example 42 was prepared above was similarly digested to produce a fragment of about 640 bp after which the two fragments were ligated PCFM536/hMKT(1-163) prepared in Example 52 was digested with Xbal and Sfil to afford an about 270-bp fragment which digested with Xbal and HindIII, an about 1040-bp fragment EP-A-136490) predigested with Xbal and HindIII (E. coli JM109 previously transformed with pMW1 (ATCC No. 39933) was employed as a host.). The resultant clone was named digested with BamHI and HindIII, and the pBL(BH)(123-332) sequence shown in SEQ ID NO:14.

The above transformant was cultured in 60 ml of medium containing 50 µg/ml of ampicillin and 12.5 µg/ml of tetracycline overnight at 30°c, and the culture (25 ml) was then added to 1000 ml of LB medium containing 50 µg/ml of ampicillin followed by shaking culture at 36 c until the OD600 reaches 1.0 to 1.2. Then, after about 330 ml of LB medium at 65.c was added to the culture so that the final temperature of the culture became 42.c, the shaking culture was continued for further 3 hours at 42.c in order to induce expression of the variant human TPO, hMKT(1-332) protein, also referred to as Met-2, Lys-1JTPO(1-332).

(Dai-ichi Chemical Co.) was employed. After electrophoresis nitrocellulose membrane), the above expressed protein was Example 45. After staining, a band was detected at a The obtained culture was directly subjected to SDS-PAGE analysis. In the SDS-PAGE analysis, Multigel 15/25 and Coomassie Blue staining, the protein specific for the induction of expression was detected at a molecular weight of about 35 KD on the gel with regard to the transformant that addition, following SDS-PAGE and electroblotting (using eacted with the anti-HT1 peptide antibody prepared in molecular weight of approximately 35 kD thus indicating that the expression of the hMKT(1-332) protein was induced. he hMKT(1-332) was expressed.

<Example 67>

their expression in COS7 cells, and identification of their Preparation of human TPO substitution derivatives.

several derivatives that the amino acids of human TPO were partially substituted with an other amino acid(s) have TPO According to the following procedures, whether activity was studied.

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The vector pSMT201 for expression in animal cells, for use in this Example, was constructed as follows.

and EcoRI, and then electrophoresed on agarose gel to recover a DXM vector fragment wherein the cDNA for murine Next, two oligonucleotides for introducing various cloning restriction First, the expression plasmid pXM-mEPORn which contains a cDNA for murine erythropoietin receptor (obtained rom Dr. D'Andrea in Dana Farbor Cancer Institute; Cell: 57, 277-285 (1989)) was treated with restriction enzymes Kpnl sites into the above expression vector were synthesized using a DNA synthesizer (ABI). The oligonucleotides synthesized erythropoietin receptor was removed. have the following basic sequences:

primer-1:

primer-2:

DOGGOGTOCATCADCATCACTGAGOGGOCGCCC3 (SEQ ID NOA58) SCCIOSAGGAATICCIGCAGCCCGGGACIAGIAICGGCTACCCCTACCACGICCCCGACIACG

SAATTESCOSCOCCICAGTEGTCATGGTGATGAGCCCCCCGGTAGTCCSGGGACGTCGTAG GESTAGOCGATACTAGTCCCGGGCTTGCAGGAATTCCTCGAGGGTAC-3 (SEQIDNO-159)

were ligated together using T4 DNA ligase (Takara-Shuzo) to produce an expression vector pSMT201. The pSMT201, as to form a double stranded oligonucleotide which was thereafter ligated with the pXM vector recovered above in the presence of T4 DNA ligase (Takara-Shuzo) to give an expression vector pDMT201. To remove the murine DHFR cDNA followed by agarose gel electrophoresis to recover a larger ragment from the pEF18S vector DNA. Then, these fragments shown in Fig. 21, has an origin of replication of SV40, an The two oligonucleotides were mixed and annealed contained in pDMT201 from the pDMT201, the vectors pDMT201 and pEF18S were separately digested with Notl and Hpal ragment from the pDMT201 vector DNA and a smaller

into a vector pSMT201 predigested similarly to produce a sequence, an SV 40 early polyadenylation site sequence, an DUC18 and a B-lactamase gene (Amp^r), together with the Pstl, Kpnl, Xhol, EcoRl, Smal, Spel and Notl sites. The pHTP llustrated in Example 30 carrying the full length human TPO numan TPO cDNA fragment which was subsequently subcloned enhancer sequence, adenovirus major late promoter sequence, an adenovirus tripatite leader sequence, a splice signal adenovirus VA RNA gene sequence, an origin of replication of ollowing restriction sites for ligation of a desired gene: BgIII, cDNA was digested with EcoRI and Spel to give a full length plasmid pSMT201-hTPO.

Using the thus obtained plasmid pSMT201-hTPO as a template, the plasmid BGL-TPO/pBlue which was employed as a template for the preparation of derivatives of interest was firstly constructed as follows.

human TPO was attempted by the Annweiler's method For this purpose, the following two DNA strands which were globin leader sequence to the 5'-non-translational site of In the BGL-TPO/pBlue the addition of the rabbit \beta-(Annweiler et al., Nucleic Acids Research, 19: 3750, 1991). chemically synthesized:

S-AATTCCAAGATCTCACACTTGCTTTTGACACAAC

TRESTITACTTCCAAATCCCCCAAAACAGACAGACCC3" (SEQ ID NO:160); and SGGGICTGTCTGTTTGGGGGGATTGCAAGTAAACA

EcoRI and Smal to produce a vector BGL/pBlue in which the digesting the vector Bluescript II SK+ (Toyo-boseki) with CAGTTGTGTCAAAAGCAAGTGTGAGATCTTGG-3 (SEQ ID NO:161), were ligated with a fragment which had been obtained by eader sequence was inserted.

PCR was conducted using the following primer sednences:

Sma-ATG: 5'-GGCCCGGGATGGAGCTGACTGAATTGCTC-3'

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(SEQ ID NO:162)

(i.e., 102-122 sequence of SEQ ID NO: 7 to which Smal sequence has been linked); and

end: 5'-TCAAGCTTACTAGTCCCTTCCTGAGACAGATTCTG-3'

(i.e., antisense of the 1140-1160 sequence of SEQ ID NO.7, to which Spel and HindIII sequences had been linked). The PCR was carried out using 1 ng of the plasmid pSMT201-hTPO DNA as a template together with 10 µM the Using TaKaRa PCR Amplification Kit Research), the PCR was conducted in a volume of 100 µl under (Takara-Shuzo) and Programmable Thermal Controller (MJ the following conditions: 3 cycles of 94 c 1 min, 55 c 2 min min and 72 c 1 min/cycle. The PCR product was extracted and 72.c 2 min/cycle; then 20 cycles of 94.c 45 sec, 55.c 1 with chloroform and then ethanol-precipitated twice followed by suspension in 100 µl of TE buffer. synthetic primers.

phenol/chloroform, and precipitated with ethanol. The Competent High E. coli JM109 (Toyo-boseki) was used as a host). Of the transformant cells obtained, twenty clones were was confirmed that the plasmid encoding the ßGL-TPO/pBlue Subsequently, the PCR product was subjected to the restriction digestion with Smal and HindIII, extracted with esultant precipitate was dissolved in 10 µl of TE buffer, after which it was subcloned into the vector BGL/pBlue which was previously digested with the same restriction enzymes Harbor Laboratory Press (1989). The purified plasmid DNA was dentified through sequencing using Taq Dye Deoxy^{T M} Terminater Cycle Sequencing Kit (Applied Biosystems, Inc.) and 373A DNA sequencer (Applied Biosystems, Inc.). As a result, it nad the expected TPO cDNA sequence without any substitution over the full length thereof. Then, after the BGL-TPO/pBlue selected from which plasmid DNA's were prepared essentially as described in Sambook et al., Molecular Cloning, Cold Spring

cDNA were linked at the BgIII/Spel restriction site vector as shown in Fig. 21. This pSMT/BGL-TPO vector was precipitate obtained was dissolved in 10 µl of TE buffer, and it structure that the β-globin leader sequence and the human TPO was digested with BgIII and Spel, the digest was extracted was subcloned into the vector pSMT201 which has been digested with the same enzymes (Compitent-High E. coli JM109 Toyo-boseki) was used as a host.). In accordance with the cells. The resulting expression vector pSMT/BGL-TPO had the nethod described in the Molecular Cloning (Sambook et al., supra), the plasmid DNA was prepared from the transformant downstream of the sprice signal sequence of the pSMT201 with phenol/chloroform and then ethanol-precipitated. amployed for its transfection into COS cells.

For the preparation of human TPO substitution derivatives, PCR was applied in which the Ito's method was two derivatives of human TPO that the Arg-25 and His-33 of were illustratively prepared. Respectively, these derivaticves can be referred to as [Asn²⁵]TPO and [Thr³³]TPO. In PCR the employed (Ito et al., Gene, 102:67-70, 1991). In particular, the human TPO are substituted by Asn and Thr respectively ollowing primers were employed:

(corresponding to T7 promoter region of Bluescript II SK+); ABBIII: 5'-AATTCCAAGATCACACATTGC-3' (SEQ ID NO:165) and: 5'-TCAAGCTTACTAGTCCCTTCCTGAGACAGATTCTG-3' 17: 5'-TAATACGACTCACTATAGGGCG-3' (SEQ ID NO:164) (for substitution of BgIII recognition sequence);

(antisense of the 1140-1160 of SEQ ID NO:7, to which Spel and (SEQ ID NO:166) Hindlil sequences have been linked);

VB: 5-TGGGCACTGGCTCAGGTTGCTGTGAAGGACATGGG-3"

(SEQ ID NO:167)

(Arg 25 -> Asn of SEQ ID NO:7); and

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The first step PCR was carried out using 1 ng of the plasmid BGL-TPO/pBlue DNA as a template together with 10 µg each of he synthetic primers. The combinations of the primers were: Jsing TaKaRa PCR Amplification Kit (Takara-Shuzo) and Programmable Thermal Controller (MJ Research), the PCR was conditions: 3 cycles of 94 c 1 min, 55 c 2 min and 72 c 2 min/cycle; then 17 cycles of 94 c 45 sec, 55 c 1 min and 72 c min/cycle. Each of the PCR products was extracted with chloroform and precipitated twice with ethanol, and the precipitate was dissolved in 100 µl of TE buffer. The egard to, the primers, the combination of T7 and end was 10 µl of TE buffer, and subsequently subcloned into the conducted in a volume of 100 µl under the following resulting PCR products (1 µl each) were subsequently subjected to a second PCR. The combinations of the templates were: PCR products of [1]/[2] and PCR products of [1]/[3]. With employed in two cases. After incubation at 94.c for 10 min and addition of TaKaRa Taq, the PCR was carried out under the ollowing conditions: 3 cycles of 94 c 1 min, 55 c 2 min and 72.c 2 min/cycle; then 9 cycles of 94.c 45 sec, 55 ·c 1 min and 72 c 1 min/cycle. To each of the resulting PCR products were added 1 μl of proteinase K (5 mg/ml), 2 μl of 0.5 M EDTA and 2 µl of 20% SDS, incubated at 37°c for 30 min to nactivate Taq, extracted with phenol/chloroform, and ethanol-precipitated. Thereafter, the precipitate dissolved in 20 µl of sterile water was digested with BgIII and Spel, precipitation. The precipitate thus obtained was dissolved in expression vector pSMT201 which has been previously digested [1] ABgill and "end" primers; [2] T7 and N3; and [3] T7 and O9. ollowed by phenol/chloroform extraction and ethanolwith the same restriction enzymes and treated with calf

the transformant cells obtained, in each case two clones were essentially as described in the Molecular Cloning (Sambook et al., supra). The purified plasmid DNAs were sequenced using 373A DNA sequencer and Taq Dye DeoxyTM Terminater Cycle (Competent-High E. coli: JM109 was employed as a host.). Of selected from which the plasmid DNAs were prepared intestine alkaline phosphatase (Boehringer-Mannheim) Sequencing Kit (both, Applied Biosystems, Inc.).

primers [1] and [3] contained a cDNA encoding a TPO The plasmid prepared by PCR with the use of the substitution derivative (09/TPO). The amino acid substitution of His33(CAC) by Thr(ACC) and the extension of the original C-This derivative was referred to as [Thr33, Thr333, Ser334, IIe335, Gly336, Tyr337, Pro338, Tyr339, Asp340, Val341, Pro342, Asp343, Tyr344, Ala345, Gly346, Val347, His348, terminus (Gly 332) by the addition of the amino acid sequence, TSIGYPYDVPDYAGVHHHHH (SEQ ID NO:169), were confirmed. His 349, His 350, His 351, His 352, His 353 TPO.

TSIGYPYDVPDYAGVHHHHHH, were confirmed. This derivative was referred to as [Asn²⁵, Lys²³¹, Thr³³³, Ser³³⁴, Ile³³⁵, Gly³³⁶, Tyr³³⁷, Pro³³⁸, Tyr³³⁹, Asp³⁴⁰, Val³⁴¹, Pro³⁴², Asp³⁴³, Tyr³⁴⁴, Ala³⁴⁵, Gly³⁴⁶, Val³⁴⁷, His³⁴⁸, His³⁴⁹, encoding a TPO substitution derivative (N3/TPO). The amino acid substitutions of Arg25(AGA) by Asn(AAC) and On the other hand, the plasmid prepared by PCR with the use of the primers [1] and [2] contained a cDNA Glu231(GAA) by Lys (AAA) and the extension of the original Cterminus(Gly332) by the addition of the amino acid sequence, His 350, His 351, His 352, His 353 TPO.

The transfection into COS7 cells of each clone obtained was carried out by the DEAE-dextran method including chloroquine treatment as described in Example 35. Briefly, 40 after 5 days, the culture supernatant was recovered which ug each of the plasmid DNA's were used in transfection, and,

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COS7 cells into which plasmids containing cDNAs encoding the numan TPO substitution derivatives N3/TPO and O9/TPO were ransfected respectively, the TPO activity was detected in a were subsequently concentrated to a volume of 1/20 using by the M-O7e assay. As a result, in the culture supernatants of Centricon-30 (Amicon) in order to evaluate their TPO activity dose-dependent manner (see Fig. 22).

<Example 68>

expression thereof in COS7 cells and identification of TPO Preparation of insertion or deletion derivatives of human TPO. activity This Example makes evident whether insertion or deletion derivatives of hTPO163 have a TPO activity. Derivatives prepared were a His33-deletion derivative ([AArg¹¹⁷]TPO(1-163), dR116); a Thr-insertion derivative ([His³³, Thr³³', Pro³⁴]TPO(1-163), T33'), an Ala-N116'), an Ala-insertion derivative ([Gly¹¹⁶, Ala^{116'}, Arg¹¹⁷]TPO(1-163), A116') and a Gly-insertion derivative ([ΔHis^{33}]TPO(1-163), dH33); a Gly116-deletion derivative ([ΔGly¹¹⁶]TPO(1-163), dG116); an Arg117-deletion Ser38jTPO(1-163), G33") in which Thr, Ala and Gly have been Ala and Gly have been independently inserted between Gly116 insertion derivative ([His³³, Ala³³', Pro³⁴]TPO(1-163), A33') ndependently inserted between His33 and Pro34; and an Asn-[Gly¹¹⁶, Gly¹¹⁶', Arg¹¹⁷]TPO(1-163), G116') in which Asn, and a Gly-insertion derivative ([His³³, Gly³³', Pro³⁴ insertion derivative ([Gly¹¹⁶, Asn¹¹⁶', Arg¹¹⁷]TPO(1-163) and Arg117, all the sequences being based on SEQ ID NO:13. derivative

Among the derivatives, T33' and N116' are intended to introduce mucin type and Asn-binding type sugar chains,

The above mentioned derivatives were prepared using och by the method described by Ito et al (Gene, 102: 67-70, 1991). Primer sequences used in PCR are as follows:

for the preparation of His33-deletion derivative based on the JH33: 5'-TGTAGGCAAAGGAACCTCTGGGCA-3' (SEQ ID NO:172) sequence shown in SEQ ID NO:7);

133: 5'-TGTAGGCAAAGGAGTGTGAACCTCTGG-3' (SEQ ID NO:173) (for the preparation of Thr-insertion derivative with Thr inserted between His33 and Pro34 in the sequence shown n SEQ ID NO:7);

433: 5'-TGTAGGCAAAGGAGCGTGAACCTCTGG-3' (SEQ ID NO:174) (for the preparation of Ala-insertion derivative with Ala inserted between His33 and Pro34 in the sequence shown in SEQ ID NQ:7);

933: 5-TGTAGGCAAAGGTCCGTGAACCTCTGG-3' (SEQ ID NO:175) (for the preparation of Gly-insertion derivative with Gly inserted between His33 and Pro34 in the sequence shown n SEQ ID NO:7);

dG116: 5'-AGCTGTGGTCCTCTGTGGAGGAG-3' (SEQ ID NO:176) for the preparation of Gly116-deletion derivative based on the sequence shown in SEQ ID NO:7);

JR117: 5-GTGAGCTGTGGTGCCCTGTGGAGG-3' (SEQ ID NO:177) for the preparation of Arg117-deletion derivative based on he sequence shown in SEQ ID NO:7);

Asn inserted between Gly116 and Arg117 in the sequence N116: 5'-AGCTGTGGTCCTGTTGCCCTGTGGAGG-3' (SEQ ID VO:178) (for the preparation of Asn-insertion derivative with shown in SEQ ID NO:7);

4116: 5'-AGCTGTGGTCCTAGCGCCCTGTGGAGG-3' (SEQ ID Vla inserted between Gly116 and Arg 117 in the sequence VO:179) (for the preparation of Ala-insertion derivative with thown in SEQ ID NO:7);

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for the substitution of a restriction enzyme EcoRI-recognition Jari: 5'-cgggctgcaggatatccaagatctca-3' (SEQ ID NO:181) sedneuce);

andNotl: 5-GGGGGGCGCTCAGCTGGGGACAGCTGTGGTGGG-3' SEQ ID NO:183) (antisense of the 633-653 nucleotide sequence of SEQ ID NO:7, to which a termination codon and a Notl 77: 5'-TAATACGACTCACTATAGGGCG:3' (SEQ ID NO:182) (corresponding to T7 promoter region of BluescriptlISK+); and recognition sequence have been added).

A first step PCR was carried out using, as a template, 1 ng of the plasmid BGL-TPO/pBlue DNA prepared in Example 67 and using 10 µM synthetic primers. The and mutated primers (dH33, A33', G33', T33', dG116, dR117, A116', G116' and N116'). In PCR, TaKaRa PCR Amplification Kit Research) were employed, and the PCR was carried out in a linal volume of 100 µl. In the first PCR reaction of [1], one cycle was 94 c 1 min, 45 c 2 min and 72 c 2 min and 3 cycles were conducted, followed by 17 cycles in which one cycle was combination of the primers was: [1] dRI and endNotl or [2] T7 Takara-Shuzo) and Programmable Thermal Controller (MJ the first PCR reaction of [2] was carried out under the conditions of 94.c 1 min, 55.c 2 min and 72.c 2 min for 3 cycles followed by 94.c 45 sec, 55.c 1 min and 72.c 1 min for 17 cycles. Each of the PCR products was extracted with chloroform before twice ethanol precipitation, and then the 94.c 45 sec, 45.c 1 min and 72.c 1 min. On the other hand, esultant precipitate was dissolved in 100 µl of TE buffer.

Next, a second step of the PCR was conducted using 1 ul of each of the PCR products of [1] and [2]. The primers used were a combination of T7 and endNotl. After incubation at

TakaRa Taq. The second PCR was carried out under the conditions of 94.c 1 min, 55.c 2 min and 72.c 2 min for 3 shenol/chloroform followed by ethanol-precipitation, and the esultant precipitate was then dissolved in 20 µl of sterile water. After digested with EcoRI and Notl, the solution was ntestine, Boehringer-Mannheim). A host cell employed was ransformant thus obtained, a plasmid DNA was prepared essentially by the method described in Molecular Cloning Biosystems 373A DNA sequencer. As a result, it was cDNA sequence without any substitution of a nucleotide 34.c for 10 min, to each PCR reaction mixture was added cycles followed by 94.c 45 sec, 55.c 1 min and 72.c 1 min for 3 cycles. To each PCR product thus obtained were added 1 µl of 5 mg/ml proteinase K, 2 µl of 0.5 M EDTA and 2 µl of 20% SDS, and the mixture was kept at a temperature of 37°c for 30 min to inactivate Taq. The PCR product was extracted with extracted with phenol/chloroform and then precipitated with ethanol. The obtained precipitate was dissolved in 10 µl of TE buffer, and subcloned into an expression vector pEF18S that nas been previously digested with the same restriction enzymes and treated with alkaline phosphatase (from calf Competent-High E. coli JM109 (Toyo-Boseki). From the Thereafter, the nucleotide sequence of the purified plasmid DNA was confirmed using Taq Dye DeoxyTM Terminater Cycle Sequencing Kit (Applied Biosystems) and using an Applied confirmed that all the plasmids encoding dH33, A33', T33', dG116, dR117, A116', G116' and N116' had the expected TPO sequence at sites other than the intended sites. Moreover, in G33' a base substitution [Pro38 (CCT)->Ser (TCT)] other than (Sambook et al., Cold Spring Harbor Laboratory Press (1989)). hat at the intended site was observed.

cells was conducted in accordance with the method in Example Briefly, the transfection was carried out using 10 µg of The transfection of each clone obtained into COS7

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each plasmid DNA by the DEAE-dextran method including chloroquine treatment, and after 4 to 5 days, the culture supernatant was recovered which was then evaluated by M-07e cell cultures transfected with plasmids encoding an amino acid insertion or deletion derivative, the TPO activity was detected in a dose-dependent manner (see Fig. 23). On the contrary, no TPO activity was found in the supernatant of the COS7 cell culture that the expression plasmid pEF18S not containing a cDNA encoding a TPO derivative has been assay system. As a result, in each supernatant of the COS7 ransfected into and expressed.

<Example 69>

Preparation and purification of anti-human TPO peptide antibodies

considered relatively suitable as antigens were selected from which a four-stranded peptide of a multiple antigen peptide Six regions (shown in Table 8) which are the amino acid sequence of human TPO of SEQ ID NO: 191, from Sci., USA, 85, 5409-5413, 1988). Two rabbits were immunized (MAP) type was synthesized by Tam's method (Proc. Natl. Acad. eight times each with 100 µg of this peptide. Ξ

Apart from the above, single-stranded peptides were produced by bonding cysteine residue to the C-terminal of each of the peptide regions shown by SEQ ID NOS: 119-124 respectively, which are partial peptides corresponding to 8 to 28, 47 to 62, 108 to 126, 172 to 190, 262 to 284, and 306 to these as test antigens, the antibody values thereto in the sera value was confirmed in all the sera tested. Thus, these sera these are referred to as peptide region HT 1 to 6 regions, 332, respectively, of the sequence of SEQ ID NO: 191). Using obtained from the thus-immunized rabbits were measured by enzyme-immunoassay, from which the increase in the antibody were referred to as anti-sera.

In addition, the above-mentioned single-stranded synthetic peptide having the cysteine residue bonded to its Cterminal was used also as the antigen for the affinity purification of the anti-sera in the following (2).

derived from each of all the thus-immunized rabbits was prepared separately. Concretely, for anti-HT1 peptide anti-HT1-2 peptide antibody derived from the two immunized (2) Two rabbits each were immunized with one of the above-mentioned antigen peptides, and the antibody antibodies, obtained were anti-HT1-1 peptide antibody and rabbits.

As one example, the purification of anti-HT1-1 peptide antibody is illustrated hereunder.

with a coupling buffer (50 mM Tris, 5 mM EDTA-Na, pH 8.5) of 6 mentioned coupling operation was conducted at room septide was coupled onto the get by covalent bonding, at a coupling efficiency of 28.3 %, to prepare an antigen peptide gel having 0.8 mg of the peptide coupled onto 1 ml of the gel, in he antigen column. After collecting all blood from each minutes, the gel was washed with the coupling buffer of 3 stand as it was for 30 minutes, the gel was washed with the coupling buffer of 8 times by volume the gel. The aboveemperature. In this way, the antigen region-containing having cysteine residue bonded thereto were coupled onto 12 ml of Sulfo Link Coupling Gel (produced by Pierce Co., as catalog No. 44895). Precisely, a peptide solution containing the antigen was coupled onto the gel that had been equilibrated imes by volume the gel, over a period of 15 minutes. Next, after having been left to statically stand as it was for 30 limes by volume the gel. Next, the coupling buffer containing 0.05 M L-cysteine-HCl was added to the gel at a rate of 1 ml/ml-gel, by which the non-treated groups were blocked over a period of 15 minutes. After having been left to statically First, 30 mg of the single-stranded peptide of HT1

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mM of NaCl and 0.05 % sodium azide. Thus, 11.2 ml of a mmunized rabbit, 78.4 ml of anti-serum containing anti-HT1l peptide antibody were obtained. 76.7 ml of the anti-serum having a protein content of 3620 mg) were added to the antigen column that had previously been equilibrated with 50 mM phosphate buffer (pH 8) containing 150 mM of NaCl and 0.05 % of sodium azide, then the column was washed with the same content of 3680 mg) were obtained. Next, the fraction which was then immediately neutralized by adding thereto 21.1 ml of 0.1 M carbonate buffer (pH 9.9), concentrated by and purified in 50 mM phosphate buffer (pH 8) containing 150 purified anti-HT1-1 peptide antibody (having a protein content as above, anti-HT1-2 peptide antibody (60.0 mg), anti-HT2-1 buffer. Thus, 105.9 ml of a passing fraction (having a protein adsorbed was eluted with 0.1 M citric acid buffer (pH 3.0), ultrafiltration (using YM30 membrane produced by Amicon Co.) In the same manner septide antibody (18.8 mg), anti-HT2-2 peptide antibody (8.2 of 77.7 mg) were obtained in the buffer. mg), etc. were obtained.

Anti-HT3 to anti-HT6 peptide antibodies are obtained in the same manner as above.

<Example 70>

Detection of TPO Western Analysis

(1) To evaluate the anti-sera obtained in Example 59, these were tested in the manner mentioned below.

had been partially purified from the supernatant of the culture to 332 of SEQ ID NO:6 had been introduced and expressed, was according to an ordinary method and then electrically blotted on a PVDF or nitrocellulose membrane. After the blotting, the A standard sample of recombinant human TPO that membrane was washed with 20 mM Tris-HCl and 0.5 M NaCl (pH of CHO cells, into which a gene coding for the amino acids -21 subjected to SDS-polyacrylamide electrophoresis (SDS-PAGE)

each with TTBS for 5 minutes. Next, this was treated with a 1/5000 dilution of alkaline phosphatase-labeled avidin 5 minutes and then with TBS for 5 minutes. Then, this was which verified that the human TPO was recognized and oeptide antibody, anti-HT4-1 peptide antibody, anti-HT5-2 diluted with TTBS solution containing 0.05 % BSA and 10 % Precisely, the recombinant human TPO sample processed as above was treated with TTBS solution containing the anti-TPO minutes and then washed with TTBS for 5 minutes. Next, this was treated with TTBS solution containing anti-rabbit(ab') L43015), as the secondary antibody, along with 0.05 % BSA and 10 % Block Ace, for 60 minutes, and then washed two times diluted with 10 % Block Ace-containing TTBS solution, for 30 minutes, and thereafter washed two times each with TTBS for developed, using an alkaline phosphatase substrate (produced by Bio-Rad Co., as catalog No. 170-6432). The abovementioned Western assay was conducted at room temperature, % Tween 20-containing TBS (TTBS) for 5 minutes, and treated with a blocking agent (Block Ace; produced by Dal-Nippon peptide antibody, anti-HT2-1 peptide antibody, anti-HT3-2 oeptide antibody and anti-HT6-1 peptide antibody each were Block Ace to 1/1000 dilutions. The thus-diluted antibodies peptide antibody, 0.05 % BSA and 10 % Block Ace for 60 goat biotinated antibody (produced by Caltag Co., as catalog No. The anti-sera each separately containing one of anti-HT1-1 were employed for Western assay as the primary antibodies. (produced by Leinco Technologies Co., as catalog No. A108) 7.5) (TBS) for 5 minutes, then washed two times each with 0.1 Pharmaceutical Co., as catalog No. UK-B25) for 60 minutes. detected by the anti-sera.

anti-HT1-2 peptide antibody, anti-HT2-1 peptide antibody and anti-HT2-2 peptide antibody that had been purified by affinity chromatography in Example 69 were weighed and biotinated by (2) 3 mg of each of anti-HT1-1 peptide antibody,

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(1) A solution comprising 50 mM Na phosphate and prepared. 0.8 ml of this solution were combined and was recovered from the reaction mixture by centrifugation. 10 ml of pure water were added thereto and, only the gel part was recovered from this by centrifugation. This process was repeated four times, by which the non-reacted antibody was 7.0) and 1.1 ml of the solution of reducing agent that had been added thereto, at 4 c for 2 hours, by which the active groups in the non-reacted gel were blocked. Last, the resulting gel was which was washed with 3 M sodium thiocyanate solution and 0.15 M NaCl (pH 8.0) and containing 5 mg/ml of the antibody coupled with 1.54 ml of a swollen formyl-activated gel Formyl-Gellulofine, produced by Chisso Co.) at 4 c for 2 hours. Vext, 1.1 ml of a solution containing 10 mg/liter of a reducing agent (trimethylamine borane (TMAB), produced by Seikagaku Kogyo K.K., as catalog No. 680246) were added thereto and coupled with them for further 4 hours. Then, only the gel part emoved. Next, the thus-recovered gel was treated with 2.1 ml of a blocking buffer (0.2 M Na phosphate, 1 M ethanolamine, pH using a centrifuge. This was filled in a small column tube, washed with water, 20 mM Tris-HCl and 0.15 M NaCl (pH 8.0),

The anti-TPO peptide antibody gel in the anti-HT1each IgG fraction. The amount of the IgG fraction coupled onto the gel per the unit volume of the gel was 1.9 mg/ml-gel. In he same manner as above, a gel was prepared, onto which 21.8 2 antibody column had a coupling efficiency of up to 94.2 % for mg, per ml of the gel, of IgG not having TPO as the antigen had been coupled. This was used as a pre-column (hereinafter referred to as a pre-antibody column) for the purpose of emoving the non-specifically-bonding molecules from TPO intibody columns, as is mentioned in (2), equilibrated column was stored.

0.1 M glycine-HCl (pH 2.5) solution and then again equilibrated

with 20 mM Tris-HCl and 0.15 M NaCl (pH 8.0).

The thus-

standard sample of recombinant human TPO as that used in the coupling them onto an activated biotin (NHS-LC-Biotin II, produced by Pierce Co., as catalog No. 21336). The same oregoing (1) was subjected to SDS-PAGE, then electrically olotted on a PVDF or nitrocellulose membrane, and subjected to ordinary Western assay using each of these biotinated antibodies as the primary antibodies. Immediately after the plotting, the membrane was washed with TBS for 5 minutes and then washed two times each with TTBS for 5 minutes, and or 60 minutes. Next, this was treated with TTBS solution containing 1 µg/ml of the biotinated anti-TPO peptide antibody, 0.05 % BSA and 10 % Block Ace for 60 minutes and his was treated with a 1/5000 dilution of alkaline phosphatase-labeled avidin (produced by Leinco Technologies thereafter this was treated with a blocking agent (Block Ace) Co., as catalog No. A108) diluted with 10 % Block Acecontaining TTBS solution, for 30 minutes, and thereafter IBS for 5 minutes. Then, this was developed, using an alkaline The above-mentioned Western assay was then washed two times each with TTBS for 5 minutes. Next, washed two times each with TTBS for 5 minutes and then with conducted at room temperature, which verified that the human phosphatase substrate (produced by Bio-Rad Co., as catalog No. TPO was recognized and detected by the purified antibodies. 170-6432).

Preparation of anti-TPO peptide <Example 71>

The anti-human TPO peptide antibodies obtained in Example 69 were verified to recognize human TPO. These antibody columns and detection of human TPO

antibodies were separately coupled onto a chromatography carrier to prepare anti-TPO peptide antibody columns. As one example, the preparation of anti-HT1-2 peptide antibody

column is illustrated hereinafter.

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(2) Next, another example of preparing an anti-HT1-2 antibody column is illustrated hereinafter.

2.5). Next, the fraction that had passed through the prewere passed through the column, whereupon the fraction antibody column was added to the anti-HT1-2 antibody column (containing 2 ml of gel) prepared in (1), and the column was through the anti-HT1-2 antibody column. Next, the fraction adsorbed to the anti-HT1-2 antibody column was eluted with 8 and that almost all TPO in the sample added bonded the latter column. From this, it was confirmed that at least from 200 to had been partially purified from the supernatant of the culture of CHO cells into which a gene coding for the amino acid sequence of any of SEQ ID NOS: 119-124 had been introduced and expressed was added to the pre-antibody column volume the gel of 20 mM Tris-HCl (pH 8.0) and 0.15 M NaCl passing through the column was collected. Next, the fraction washed with 10 times by volume the gel of 20 mM Tris-HCl (pH 8.0) and 0.15 M NaCl, while collecting the fraction passing OH 2.5). These fractions were analyzed by SDS-PAGE, which verified that TPO did not adsorb to the pre-antibody column out was specifically bonded to the anti-HT1-2 antibody column 300 µg/ml-gel of TPO was bonded to the gel in the latter A standard sample of recombinant human TPO that containing 1 ml of gel) prepared in (1), and 10 times by adsorbed to the pre-antibody column was eluted with 10 times by volume the gel of an acidic eluent (0.1 M glycine-HCl, pH times by volume the get of an acidic eluent (0.1 M glycine-HCl,

<Example 72>

Effect of TPO for Increasing the Numbers of Platelets

Twenty normal ICR strain male mice, eight weeks of age, (blood was collected from their orbit veins already and the numbers of platelets was measured using a micro cell

the TPO-iv group, a 177% increase was noted on Day 6 as was counted using a micro cell counter (F-890; manufactured by Toa Iyo Denshi). The changes in the numbers of platelets Day 8 where the numbers were highest. On the other hand, in in the TPO-sc group, an increase of 347% was noted already on groups (TPO-iv group) was injected with 100 microliters of a microliters of PBS subcutaneously once daily for five was injected with 100 microliters once daily for five consecutive days in the same manner as in the TPO-iv group subcutaneously. After 6, 8, 10, 13 and 15 days from the initiation of the administrations, blood was collected from the orbit yein of each of the mice and the numbers of platelets are given in Fig. 24. Thus, in the control-iv group, the platelet number increased by 44% as compared with before the administration even on the Day 6 (Day 0 being the day of in the control-sc group, only a 47% increase was noted even on compared with the numbers of prior to the administration and, Day 6 and, on Day 8, the highest increase of 493% was counter F-800 manufactured by Toa Iyo Denshi) were randomly divided into four groups. One of the four groups (a control-iv group) was injected with 100 microliters of PBS intravenously route once daily for five consecutive days. One of the other solution (prepared by substituting the concentrated solution of the TPO active fraction F2 of the SP Sepharose Fast Flow obtained in Example 56 with PSB using an NAP-25 column Pharmacia Biotech; Catalog No. 17-0852-02] followed by diluting with PBS;containing relative activity, 211,900 by M-07e assay) intravenously once daily for five consecutive days. Another group (control-sc group) was injected with 100 consecutive days while the remaining group (TPO-sc group) administration) where the platelet numbers were highest and,

From the above results, it has been confirmed that human TPO increases the number of platelets independently of

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ō in the type and also of the route the difference administration.

cExample 73>

Effect of TPO for Increasing the Number of Platelets

weeks of age, (the blood was already collected from their orbit veins and the number of platelets were measured using a a control group) was injected with 100 microliters of PBS subcutaneously once daily for five consecutive days. Another nicroliters of a solution (prepared by substituting the TPO he M-07e assay) subcutaneously once daily for five Sixteen normal C3H/HeJ strain male mice, 11 were randomly divided into four groups. One of the four groups group (TPO-1 group) was injected subcutaneously with 100 with PBS using NAP-25 followed by diluting with PBS; containing relative activity, 83,000 by the M-07e assay) once daily for five consecutive days. Still another group (TPO-2 group) was injected with 100 microliters of a solution group with PBS to an extent of 2-fold; containing relative activity, 41,500 by the M-07e assay) subcutaneously once extent of 2-fold more; containing relative activity, 20,750 by microcell counter type F-800 manufactured by Toa Iyo Denshi) active fraction of Sepha cryl S-200HR obtained in Example 56 prepared by diluting the TPO active fraction used in the TPO-1 was injected with 100 microliters of a solution (prepared by diluting the TPO active fraction used in the TPO-2 group to an taily for five consecutive days. The last group (TPO-3 group) consecutive days.

Blood was collected from the orbit veins of the nice on Day 6, 8, 10 and 12 after injection and the numbers of platelets were counted using a microcell counter (type F-800; manufactured by Toa Iyo Denshi). The changes in the numbers of platelets are given Thus, in the control group, there was nearly no Fig. 25.

which TPO was injected, there were increases in the numbers change in the numbers of platelets while, in the groups to of platelets in all of the groups to the extent that the numbers Thus, in the TPO-1 group, about a 200-% increase was noted on Day 6 already, the increase rose to about 270% on Day $oldsymbol{\beta}$ and then, however, a decrease was noted, even on Day 12, about a 65-% increase was still noted whereupon there was a significant difference from the control group (p < 0.01; by a Dunnet multiple comparison test). In the TPO-2 group, the same increase was found and, though the increasing action was smaller than the TPO-1 group, increases of about 140% Day 12, the numbers decreased to nearly the same level as the control group. The increasing action of the TPO-3 group was weaker than the TPO-2 group and, on Day 8 where the numbers were highest, about a 110-% increase was noted and, again, there was a significant difference (p < 0.01) from the ϕ ontrol were highest on Day 8 after the injection. In addition, differences in dose-response were noted among the groups. and about 160% were noted on Days 6 and 8, respectively. group.

peen confirmed that the human TPO increases the platelet numbers independently of the strain of the mice and that its action has From the above-mentioned results, it has a dose-responding property.

<Example 74>

An Inhibiting Effect of TPO on the Thrombocytopenia Caused by Administration of Anti-Cancer Agent

Thirty ICR male mice, eight weeks of age, were divided into two groups (each group comprising 15 mice). From he next day (Day 1), one of the groups (control group) was njected with 200 mg/kg of 5-FU intravenously and randomly njected with 100 microliters of PBS subcutaneously once laily for five consecutive days while another group/ (TPO

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consecutive days while another group (TPO group) was injected with 200 microliters of the TPO active fraction used in the M-07e assay) subcutaneously twice daily for consecutive with 200 microliters of PBS subcutaneously twice daily for Example 73 (without diluting with PBS) to which 0.04% of fween 80 was added (containing relative activity, 380,000 by

were randomly divided into three groups - from one group 800; manufactured by Toa Iyo Denshi). The changes in the in the TPO group, a decrease to about 38% of the value before situation changed to an increase. Thus, on Day 8, the value administration and, on Day 10 and thereafter, the numbers of After the injection, the mice in each of the group blood was collected on Days 5 and 12; from another group on Days 8 and 14; and from the remaining group on Day 10. Blood was collected from the orbit veins and the numbers of platelet number were counted by a microcell counter (type Fnumbers of platelets are given in Fig. 27. Thus, in the control group, the numbers of platelets after administration of ACNU decreased as the days passed and, on Days 8-10, the value was owest (about 29% of the value before the ACNU administration) and the recovery was only about 49% and the ACNU administration was noted on Day 5 and then the recovered to about 63% of the value before the ACNU platelets were larger than that before the ACNU administration. Thus, on Days 12 and 14, increases of up to about 74% on Days 12 and 14, respectively. On the other hand, about 300% and about 400% were noted, respectively.

As noted hereinabove, on Days 8-10 of which the decrease was noted in the control group, an increase was CNU. Accordingly, it has been confirmed that human TPO observed in the TPO group already and, on Day 10, the platelet numbers were more than those before the administration of

ive consecutive days. On Days 4, 6 and 8, each five mice were group) was injected with 100 microliters of TPO active 211,900 by the M-07e assay) subcutaneously once daily for andomly chosen from each of the groups, then blood was collected from the orbit veins and the numbers of platelet manufactured by Toa Iyo Denshi). The changes in the numbers owest (about 28% of the value prior to the administration of noted as a reaction of the decrease. Even in the case of a TPO FU decreased as the days passed and, on Day 6, the value was raction used in Example 72 (containing relative activity, were counted by a microcell counter (type E-2500; of platelets are given in Fig. 26. Thus, in the control group, the numbers of platelets after the administration of 5-FU decreased as the days passed and, on Day 6, the value was 5-FU) though, on Day 8, an approximately 1.3-fold increase was of 5-FU). However, the difference between the numbers of platelets on Days 4 and 6 was very small and, on Day 6, a high value as compared with the control group was group, the numbers of platelets after the administration of 5the lowest (about 52% of the value before the administration significantly (p < 0.05; by a Dunnet multiple comparison test) maintained. On Day 8, there was an increase of about 200% of the value before the 5-FU administration.

From the above-mentioned result, it has been confirmed that human TPO inhibits the decrease of the platelet number as caused by the anti-cancer agent.

<Example 75>

A Therapeutic Effect of TPO for Thrombocytopenia

njected intravenously to each of 36 ICR male mice, seven weeks of age, and then the mice were randomly divided into wo groups (each group comprising 18 mice). As from the next Nimustine hydrochloride (ACNU) (50 mg/kg) was day (Day 1), one of the groups (control group) was injected

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promotes the recovery from the thrombocytopenia caused by he anti-cancer agent.

When the results of Example 74 are taken into consideration as well, it is now expected that human TPO inhibits the decrease in the numbers of platelets and promotes the recovery from thrombocytopenia independently of the type of the anti-cancer agent.

cExample 76>

A Therapeutic Effect of TPO on Thrombocytopenia after BMI

Forty-eight male mice of C3H/HeN strain, seven weeks of age, were irradiated with 10 Gy of radioactive were intravenously injected with 1 x 106 of bone marrow cells from the mice of the same strain. The mice were then while another group (TPO group) was injected with the TPO active fraction used in Example 73 (containing relative activity, 44,000 by the M-07e assay) with a dose of 100 microliters each subcutaneously once daily for twenty radiation on the entire body and, immediately thereafter, they andomly divided into two groups and, from the next day (Day I), one of the groups (control group) was injected with the PBS consecutive days.

collected from their orbit veins and the numbers of platelets were randomly selected on Days 5, 10, 14 and 21, blood was After initiation of the injection, each six mice were counted by a microcell counter (type E-2500; manufactured by Toa Iyo Denshi).

thereto and, on Day 10, the numbers were lowest (about 3% of gradual recovery was noted and, on Day 4, the numbers were about 11% and about 13% in the control group and in the TPO The changes in the numbers of platelets are given the numbers of before the BMT application). After that, a Thus, in all of the groups, the numbers of platelets decreased as the days passed after applying the BMT in Fig. 28.

in the control group while, in the TPO group, the recovery was confirmed that human TPO prepared as in Example 56 promote group, respectively. On Day 21, only a 37% recovery was noted about 65% whereupon a significant recovery promoting effect was observed. From the above-mentioned results, it has been he recovery of the numbers of platelets after application of

<Example 77>

A Therapeutic Effect of TPO for

Thrombocytopenia after Irradiation with Radioactive Bay

rradiation with x-rays and, on Day 9, the numbers were Thirty-six male mice of ICR strain, eight weeks of age, were irradiated with x-rays of 5 Gy on the entire body and another group (TPO group) was injected with a TPO active 1,440,000 by the M-07e assay) once daily for three relative activity, 360,000 by the M-07e assay was injected collected on Days 4, 11 and 21; another group on Days 7 and manufactured by Toa Iyo Denshi). The changes in the numbers of platelets are given in Fig. 29. Thus, in the control group, the numbers of platelets decreased as the days passed after smallest (about 25% of the number before the irradiation with X-rays). On days 11 and 13, the numbers recovered to about one of the groups (control group) was injected with PBS while subcutaneously once daily for seven consecutive days. After divided into three groups - a group from which blood was 13; and the remaining group on Days 9 and 15. The collection of blood was carried out from the orbit veins and the numbers 38% and about 67%, respectively and, on Day 15, they recovered randomly divided into two groups. From the next day (Day 1), raction used in Example 73 (containing relative activity, consecutive days subcutaneously and, from the next day, initiation of the injection, each of the group was randomly of platelets were counted by a microcell counter (type F-800)

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hose before irradiation with X-rays and, until Day 21, this to the values before the irradiation with x-rays. In the TPO group, the numbers decreased to about 24% of those before the rradiation with X-rays on Day 7. After that, an increase was Day 11 and thereafter, the platelet numbers were more than endency was maintained. As mentioned above, on Day 9 on which the numbers tended to decrease in the control group, the numbers of platelets in the TPO group already changed to an ncrease and, on Day 11, the numbers of platelets were more han those before the irradiation with X-rays while remaining ifteen days were needed for the numbers of platelets to ligh even thereafter. On the other hand, in the control group, From those results, it has been confirmed that human TPO produced in Example 56 shortens the time necessary to recover from the decrease in the numbers of platelets after irradiation with Xrays and that it exhibits an therapeutic effect for noted and, on Day 9, the numbers recovered to about 82%. thrombocytopenia after the irradiation with X-rays. recover from before the irradiation with X-rays.

<Example 78>

Effect of hTPO163 for Increasing the Numbers of Platelets

Fifteen healthy male mice of C3H/HeN strain from whom blood was already collected from their orbit veins and subjected to the number of platelets was measured by a microcell counter (type F-800; manufactured by Toa Iyo Denshi) were randomly divided into four groups - a control was injected subcutaneously with PBS containing 0.1% of serum from mouse once daily for five consecutive days. Groups A, B and C were injected with the TPO active fraction of SP Sepharose Fast Flow obtained in Example 65 followed by diluting with PBS containing 0.1% of mouse serum at the doses of about 40,000,000, about 8,000,000 and about group and groups A, B and C. The first group (control group)

1,600,000/kg body weight/day, respectively, in terms of the relative activity of hTPO163 by the M-07e assay for five consecutive days subcutaneously.

On Days 6, 8, 10 and 12 after the injection, blood was collected from the orbit vein of each mouse and the numbers of platelets were counted by a microcell counter type F-800; manufactured by Toa Iyo Denshi). The changes in he numbers of platelets are given in Fig. 30.

change in the numbers of platelets while, in the TPO-given Among each of the TPO-given groups, a dose-response Thus, in the control group, there was nearly no groups, increases in the numbers of platelets were noted in all cases giving the highest values on Day 8 after the injection. relationship was noted. Thus, in group A, increases of about 88% and about 100% were noted on Days 6 and 8, respectively and, even on Day 10, an increase of about 97% was maintained whereupon all data showed a significant difference from the Similar increases were noted in group B as well and, though the degree of increase was lower than that of group A, the increases of about 65% and about 84% were noted on Days 6 and 8, respectively showing a significant difference from the control group (p < 0.01 or 0.05 by a Dunnet Multiple Comparison Test). The increasing action in group C was lower than that in group B. Still it showed about 31% increase on Day mentioned results, it has been confirmed that the hTPO163 produced in Example 65 increases the numbers of platelets and control group (p < 0.01 by a Dunnet Multiple Comparison Test). 8 when the highest numbers were noted. From the abovehat its action shows a dose-response relationship.

Example 79>

A Therapeutic Effect of hTPO163 for Thrombocytopenia

One hundred male mice of C3H/HeN strain, eight weeks of age, were injected intravenously with 50 mg/kg of

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the groups were randomly divided into five groups and the blood was collected on Days 5, 8, 10, 12 and 14. The After initiation of the injection, mice in each of collection of blood was carried out from their orbit veins and the numbers of platelets were counted by a microcell counter (type E-2500; manufactured by Toa Iyo Denshi). The changes in the numbers of platelets are given in Fig. 31. Thus, in the control group, the numbers of platelets decreased as the days On the other hand, in group A, the numbers decreased to about that, however, they changed to increase and recovery to an passed after the administration of ACNU giving the lowest values (about 15-16% of that before the ACNU administration) on Days 8-10 and the extent of the recovery was only as little 25% of those before the ACNU administration on Day 8. After extent of about 34% and about 89% were noted on Days 10 and 12, respectively. On Day 14, the number was larger than that as about 28% and about 51% on Days 12 and 14, respectively, of before the ACNU administration.

As mentioned above, the smallest numbers of platelets were noted on Day 8 in all of the groups though, in he groups to which human TPO was given, the decrease was slower than the control group giving rise to the smallest numbers of platelets. In the control group, there was nearly

varied. In a group to which 50 micrograms/kg was in the groups given the human TPO, recovery in the numbers of platelets were noted in all of the groups, though their degrees administered, recovery was noted on Day 12 already giving he recovery was as small as about 51% as compared with the state before the ACNU administration even on Day 14, it is now clear that the hTPO163 produced as in Example 65 promotes the recovery from the decrease in the numbers of platelets. As such, the hTPO163 has been confirmed to have a therapeutic no recovery of the numbers of platelet even on Day 10 while, higher numbers than of those before the ACNU administration. When it is compared with the fact that, in the control group, effect on thrombocytopenia caused by anti-cancer agents.

Examples of the pharmaceutical preparations are hereinunder.

<Example 80>

. The human TPO fraction obtained in Example 56 concentrated, subjected to aseptic treatment and frozen at -20°c to give a frozen product to be used as an injectable preparation. was

<Example 81>

The human TPO fraction obtained in Example 56 was concentrated, 5 ml of it was filled in a 10-ml vial bottle using aseptic operation, freeze-dried at -20 c and the bottle was stopped with a rubber stopper to give a freeze-dried substance to be used as an injectable preparation.

<Example 82>

was concentrated, subjected to aseptic filtration and filled in The human TPO fraction obtained in Example

a 10-ml vial bottle to give a product to be used as an injectable preparation.

cExample 83>

at -20 c to give a frozen product to be used as an injectable The hTPO163 fraction obtained in Example 65 was concentrated, subjected to aseptic treatment and freeze-dried preparation.

<Example 84>

aseptic operation, freeze-dried at -20 c and the bottle was The hTPO163 fraction obtained in Example 65 was concentrated, 5 ml of it was filled in a 10-ml vial bottle using sealed with a rubber stopper to give a freeze-dried product to be used as an injectable preparation.

cExample 85>

10-ml vial bottle to give a product to be used as an injectable The hTPO163 fraction obtained in Example 65 was concentrated, subjected to aseptic filtration and filled in a preparation.

<Example 86>

frozen at -20 c to give a frozen product to be used as an was concentrated, subjected to an aseptic treatment and The human TPO fraction obtained in Example 57 injectable preparation.

<Example 87>

using aseptic operation, freeze-dried at -20°c and the bottle was concentrated, 5 ml of it was filled in a 10-ml vial bottle was sealed with a rubber stopper to give a freeze-dried The human TPO fraction obtained in Example 57 product to be used as an injectable preparation.

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<Example 88>

was concentrated, subjected to an aseptic filtration an filled in a 10-ml vial bottle to give a product to be used as an The human TPO fraction obtained in Example 57 injectable preparation.

<Example 89>

Aplastic Canine Plasma

492 (1987); Mazur, E., Basilico, D., Newton, J.L., Cohen, J.L., (1990)] except that 450 rads of total body irradiation were Heparinized aplastic canine plasma ("APK9") or normal canine plasma ("NK9") was produced as described Arriaga, M., South K., Cohen J.L. and Mazur, E.M. Blood 69: 486-Charland, C., Sohl, P.A., and Narendran, A. Blood 76: 1771-1782 Mazur, E. and South, K. Exp. Hematol. 13:1164-1172 (1985); delivered to recipients.

<Example .90>

Human Megakaryocyte Assay

Standard methods for many of the procedures described in the following examples, or suitable alternative procedures, are provided in widely recognized manuals of Molecular Cloning, Second Edition, ColdSpring Harbor Laboratory Press (1987) and in Ausubel, et al., (Eds), Current molecular biology such as, for example, Sambrook, et al., (Eds), Protocols in Molecular Biology, Green associates/Wiley interscience, New York (1990).

APK9 and fractionated APK9 were assayed for the ability to stimulate development of human megakaryocytes CD34- selected cells were obtained from peripheral blood cells as described (Hokom, M.H., Choi, E., Nichol, J.L., Hornkohl, A., Arakawa, T., and Hunt, P. Molecular Biology of Haematopiesis 3:15-31,1994) and were from CD34⁺ progenitor cells.

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> ncubated in the following culture medium: Iscove's modified Santa Ana, CA) and 10% heparinized, platelet-poor, human AB deoxyadenosine, 2-deoxyguanosine (10' µg/ml each, Sigma); numan recombinant insulin (10 µg/ml), human transferrin (300 1g/ml), Genzyme, Cambridge, MA); human recombinant (Vanguard, Inc., Neptune, NJ). Cells were incubated at 37·c for 8 days in humidified boxes in 5% CO2 in air, fixed directly to Dulbecco's medium (IMDM; GIBCO, Grand Island, NY) syruvic acid (110 µg/ml), cholesterol (7.8 µg/ml), adenosine, guanine, cytidine, uridine, thymidine, 2-deoxycytosine, 2-N); human recombinant basic fibroblast growth factor (2 epidermal growth factor (15 ng/ml), platelet-derived growth selected cells were plated at 2x10⁵ ml) culture medium, 15 µl linal volume, in wells of Terasaki-style microtiter plates counted with an inverted phase microscope at 100X megakaryocyte units/ml" where the degree to which a given he positive APK9 control for that experiment. One unit is supplemented with 1% Glutamine Pen-strep (Irvine Scientific, plasma. Also included were 2-mercaptoethanol (10⁻⁴ M), µg/ml), soybean lipids (1%, Boehringer Mannheim, Indianapolis, actor (10 ng/ml, Amgen, Inc., Thousand Oaks, CA). CD34he culture wells with 1% glutaraldehyde, and incubated with a Biodesign) and anti-GPIb (Dako, Carpinteria, CA). The immune reaction was developed with a streptavidin-betajalactosidase detection system (HistoMark, Kirgegaard and Perry). Megakaryocytes, identified by a blue color, were magnification. Results were presented as the average number of megakaryocytes per well +/- standard error of the mean SEM). In some cases, data were presented in terms of sample induced megakaryocyte development was normalized to defined as the amount of material that results in the same monoclonal antibody cocktail (anti-GPIb, anti-GPIIB, number of megakaryocytes as 1 µl of APK9 standard. Activity

was accepted as due to MPL ligand if it could be blocked with 5-10 µg/ml MPL-X (soluble Mpl receptor).

system. CD34-selected cells incubated with 10% NK9 for 8 days show a negligible number of blue-stained APK9 has been demonstrated to contain factor(s) hat stimulate human megakaryocyte development in this megakaryocytes, whereas CD34-selected cells incubated with 0% APK9 for 8 days show a very large number of blue-stained negakaryocytes.

Figure 32 shows that increasing concentrations of concentrations of Mpl-X greater than 5 µg/ml, inhibition is complete. In this experiment, CD34-selected cells were necessary for human megakaryocyte development, and implies Mpl-X added to the human megakaryocyte culture system stimulated with 5% APK9. This demonstrates that an activity which interacts with Mpl-X (presumptive Mpl ligand) is increasingly block megakaryocyte development. that the Mpl ligand is present in APK9 itself.

. It has been further demonstrated herein that the Mpl ligand activity necessary for human megakaryocyte development is present in APK9. APK9 (135 ml) was diluted 6fold into Iscove's media and applied to an Mpl-X affinity column. Unbound material (flow through was collected and concentrated to the original volume before assay. Bound material was eluted in 10 ml of 1 M NaCl, and 20% of the pool selected cells incubated in media alone did not develop into megakaryocytes. Cells incubated in 5% APK9 (same pool as applied to column) developed into 48 +/- 8 megakaryocytes per well. Cells incubated in 10% of the unbound material did not develop into megakaryocytes. Cells incubated in 10% of the elution pool developed into 120 +/- 44 megakaryocytes per Both the column load and the elution pool activities was diafilitered and concentrated 4-fold for assay. CD34- 268 -

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were substantially completely inhibited with 5 µg/ml Mpl-X in the assay.

<Example 91>

Transfection of murine or human Mpl receptor into a murine cell line

A. Murine Mpl Receptor

subcloned into an expression vector containing a Murine Sarcome virus. 5 µg of this construct and 1 µg of the electroporated into an IL-3 dependent murine cell line (32D, The full length murine Mpl receptor cDNA was transcriptional promoter derived from the LTR of Moloney selectable marker plasmid pWLNeo (Stratagene) were co-Geneticin (G418, Sigma) and 1 ng/ml murine IL-3. The antipeptide serum. One population was chosen for FACS cline 23; Greenberger et al., PNAS 80:2931-2936 (1983)). then incubated in selection media including 800 µg/ml surviving cells were then divided into pools of 2x10⁵ cells and analyzed. Six populations were tested for surface expression sorting using the same antipeptide serum as before. Singlecells clones of the parent cell line were selected by growth in the cells were maintained in 1 ng/ml murine IL-3. One of the Cells were cultured for 5 days to recover from the procedure, cultured until a population grew out which could be further of Mpl receptor by FACS analysis using a polyclonal rabbit 10% APK9 and Geneticin. After selection in APK9 for 35 days, subclones, 1A6.1, was used for this body of work.

Human Mpl Receptor

The full length human Mpl receptor sequence (Vigon, I., et al., PNAS 89: 5640-5644 (1992)) was subcloned into an expression vector containing the transcriptional promoter of Maloney Murine Sarcoma virus (same vector as with the murine receptor). Six µg of this construct and 6 µg

of an amphotrophic retroviral packaging construct (Landau, N.R., Littman, D.R., *J. Virology* 66: 5110-5113 (1992)) were transfected into 3 x 10⁶ 293 cells using a CaPO₄ mannalian transfection kit Stratagene. The same cells were retransfected after 2 days and again after 4 days. The day after the last transfection the 293 cells were cocultivated with the IL-3 dependent murine cell line (32D, clone 23; Greenberger et al., *PNAS* 80: 2931-2936 (1983)). After 24 hours, the 32D cells were rescued and banded in a BSA gradient (Path-o-cyte; Mills Inc.). Cells were expanded in 1 ng/ml murine IL-3 and then were selected for growth in 20% APK9. Cells were sorted for cell surface expression of receptor by FACS using a polyclonal rabbit antipeptide serum. These cells were subsequently used in the assays.

<Example 92>

1A6,1 assay for Mpl ligand

1A6.1 cells were washed free of culture IL-3 and replaced (1000 cells/15 µl total col/well) in Terasaki-style microtiter plates in alpha MEM (Gibco) supplemented with 10% fetal calf serum (FCS), Geneticin (800 µg/ml) and 1% pen/strep (Gibco) in 1:1 serial dilutions of test samples. After 48 hours, the number of viable cells per well was defined as that amount of activity that resulted in 200 viable cells per well. Activity was defined as due to Mpl ligand if it could be completely blocked by including 5-10 µg/ml Mpl-X in the assay. Mpl ligand activity in APK9 averaged 4400 +/- 539 units/ml of aplastic plasma. Unless otherwise indicated, units of Mpl ligand activity are defined in the 1A6.1 assay.

Assays with cells transfected with the human Mpl receptor gene were carried out in essentially the same manner as with the 1A6.1 cells.

. . .

Demonstration that Mpl-ligand is present in aplasticplasma of sera of mouse, dog, pig and human sources Example 93>

Mpl ligand is present in the aplastic plasma or sera Plasma was collected from BDF1 mice pre-irradiation and 12 days post-irradiation (500 rads). Plasma was tested in the that was substantially completely inhibitable with Mpl-X (10 1g/ml). Irradiated mouse plasma was also positive in the rradiation and 10 days post-irradiation (450 rads). Plasma vas tested in both the 1A6.1 assay and human megakaryocyte Mpl-X (10 μg/ml) in both assays. Plasma was collected from megakaryocyte assays. In both assays it displayed Mpl ligand activity (inhibitable by 10 µg/ml Mpl-X) comparable to that found in aplastic canine plasma. Sera from aplastic humans transplantation patients. The sera from 6 patients were assayed in the 1A6.1 assay where it showed an activity of 903 10 µg/ml Mpl-X). Sera from 14 aplastic patients has also been displaced substantial activity, 941 meg units/ml, which was completely inhibitable with 10 µg/ml Mpl-X. Murine IL-3 data Although this recombinant cytokine induces growth of the cell 1A6.1 assay where it demonstrated 2000 units/ml activity human megakaryocyte assay where it displayed an activity of Plasma was collected from dogs preassays. Activity was detected and completely inhibited by Plasma was tested in both the 1A6.1 assay and the human was obtained. This material was collected from bone marrow units/ml, 88% of which was due to Mpl ligand (inhibitable with tested in the human megakaryocyte assay. As a group, they pigs pre-irradiation and 10 days post-irradiation (650 rads). is included to demonstrate the specificity of the 1A6.1 assay. rom murine, canine, porcine and human sources (Table 9) 1833 units/ml.

anot ton	anob ton	\$9\$-/+0009	S9S-/+0009	£Մեսնա
0-/+0	841-/+146	114+-33	1 9-/+€06	nsmud sitsalqA
0-/+0	0-/+0	0-/+0	0-/+0	namud lamoM
10+/-10	1284+/-182	0-/+0	9811-/+9986	Aniorog oitzsigA
0-/+0	0-/+0	0-/+0	0-/+0	Normal porcine
0-/+0	821-/+76 <i>L</i>	0-/+0	6ES-/+00pp .	Aplistic canine
0-/+0	0-/+0	0-/+0	0-/+0	Mormal canine
anob ton	1833	0	. 5000	oznom pitzsiqA
0-/+0	0-/+0	0-/+0	0-/+0	Normal mouse
X-10M+	Media '	X-10M +	Media	
Meg Assay units/ml)		Cell Assay nits(ml)		Spicoles

TABLE 9

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Example 94>

The preparation of the human TPO derivatives expressed in E. coli system

stability and solubility, several TPO derivatives were designed replaces a lysine at amino acid position 59 (also referred to as eplaces a leucine at amino acid position 129 (also referred to wherein an arginine replaces a histidine at amino acid position at amino acid position 143 (also referred to as M143R), [Met-2, amino acid sequence set forth in SEQ ID NO:11, included [Met-2, Arg143JTPO(1-163) wherein an arginine replaces a methionine replaces a glycine at amino acid position 82 (also referred to (59R), [Met-2, Lys-1, Ala1, Val3, Arg115]TPO(1-163) wherein and expressed in E. coli. The derivatives generated using the an arginine replaces a glutamate at amino acid position 115 In an attempt to improve the biological activity, amino acid position 148 (also referred to as S148P), [Met-2, wherein a leucine replaces a glycine at amino acid position 133 (also referred to as H133R), [Met-2, Lys-1, Ala1, Val3, ys-1, Ala1, Vaß, Arg129JTPO(1-163) whèrein an arginine 146 (also referred to as G146L), [Met-2, Lys-1, Ala1, Val3, Pro148)TPO(1-163) wherein a proline replaces a serine at ys-1, Ala1, Val3, Arg59JTPO(1-163) wherein an arginine us L129R), [Met-2, Lys-1, Ala1, Val3, Arg133]TPO(1-163) ys-1, Ala 1, Val3, Leu82JTPO(1-163) wherein a leucine as G82L), [Met-2, Lys-1, Ala1, Val3, Leu146]TPO(1-163) also referred to as Q115R).

In making the TPO derivatives, the (1-163), described in example 42, was used as the template, and the oligo nucleotides which have the following sequences were synthesized:

L129R: S'-ATCTTCCGTTCTTTCCAGCACCT-3' (for the preparation of Arg-substitution derivative with Leu-129 substituted by Arg in the sequence shown in SEQ ID NO:11);

H133R: 5-TCTTTCCAGCGTCTGCGT-3' (for the preparation of Arg-substitution derivative with His-133 substituted by Arg in the sequence shown in SEQ ID NO:11);

M143R: 5'-CGTTTCCTGCGTCTGGTTGGC-3' (for the preparation of Arg-substitution derivative with Met-143 substituted by Arg in the sequence shown in SEQ ID NO:11);

GB2L: 5-GGCCAGCTTCTGCCGACCTGCCT-3' (for the preparation of Leu-substitution derivative with Gly-82 substituted by Leu in the sequence shown in SEQ ID NO:11);

G146L: 5-ATGCTGGTTCTGGGTTCTACCCT-3' (for the preparation of Leu-substitution derivative with Gly-146 substituted by Leu in the sequence shown in SEQ ID NO:11)

S148P: 5'-GTTGGCGGTCCGACCCTGTGCG-3' (for the preparation of Pro-substitution derivative with Ser-148 substituted by Pro in the sequence shown in SEQ ID NO:11); and K59R: 5'-GAAGAGCCGCGCTCAGGACATCC-3' (for the preparation of Arg-substitution derivative with Lys-59 substituted by Arg in the sequence shown in SEQ ID NO:11) Q115R:5'-CTGCCGCACGTGGCCGTACCAC-3' (for the preparation of Arg-substitution derivative with Gln-115 substituted by Arg in the sequence shown in SEQ ID NO:11).

Mutant plasmids were constructed using the Sculptor in vitro mutagenesis kit (Amersham). The Xba I - Hind III fragment derived from pCFM536/h6T(1-163) described in Example 42 was introduced into Bluescript II SK(-) phagemid vector (Stratagene, California USA) after digestion with Xba I and Hind III to obtain a plasmid pSKTPO, which was then transformed into E. coli JM109. The single-stranded DNA for the mutagenesis was prepared from pSKTPO using helper phage M13KO7 (Takara shuzo, Japan). Mutations were introduced into the TPO gene according to the supplier's protocols using the oligo nucleotides as listed above. The sequencing analysis was performed with DNA sequencing kit (Applied Biosystems, USA)

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according to the supplier's protocols and the mutations of TPO were confirmed.

genes were transformed into E. coil #261 in order to obtain the mutated pSKTPO were introduced into pCFM536 digested with Xba I and Hind III. The plasmids pCFM536 carrying mutated The Xba I - Hind III fragments prepared from transformant expressing the TPO derivative genes.

mutated pCFM 536 was performed according to Example 43 as previously described. The cell pellet of the transformant was trozen cell pellet (3 g) was suspended in 30mL of 20 mM Tris buffer (pH 8.5) containing 10 mM EDTA, 10 mM DTT and 1 mM The cultivation of E. coli #261 transformed with times at 1 minute (at least) intervals. The sonicated cells PMSF. The suspension was kept on ice and was sonicated 5 collected and stored in the freezer for at least 1 day. The were collected by centrifugation at 15000 rpm for 10 min.

1 mM PMSF and reduced by adding 50 mg of DTT. After stirring buffer (pH 8.7) containing 8 M guanidium chloride, 5 mM EDTA, The pellet was suspended in 30 mL of 10 mM Tris using diluted HCl and was then cooled to 4.c before dilution. or 1-1.5 hour, the pH of the solution was adjusted to 5.0 by

allowing the sample to stir for at least 2 days, the precipitate The solution was filtered with 2 and 1 M of urea, and equilibrated with 10 mM phosphate buffer pH 7.2) containing 15 % glycerol. The protein was eluted from Fast Flow (Pharmacia). The column was washed with 400 mL The sample solution was gradually diluted in 1.5 L was removed by centrifugation at 8000 rpm for 45 min. The of 10 mM CAPS buffer (pH 10.5) containing 30% glycerol, 3 M and then loaded to the column (2.6 x 10 cm) of CM-Sepharose sheets of #2 filter paper (a=90mm, Toyo filter paper, Japan) of 10 mM phosphate buffer (pH 6.8) containing 15 % glycerol urea, 3 mM cystamine and 1 mM L-cysteine overnight. After pH of the solution was adjusted to 6.8 with 6 M phosphoric acid, and diluted two times.

protein elution was monitored by the absorbance at 280 nm and phosphate buffer (pH 7.2) containing 15 % glycerol to the same the fraction containing TPO derivative was confirmed by SDSbuffer containing 0.5 M NaCl at a flow rate of 1.0 mL/min. the column using a linear gradient system from 10 mM PAGE and collected.

The protein elution was performed by using a linear gradient from 0.05 % TFA in H2O to 2-propanol containing 30 % CH3CN Under this conditions, the TPO derivatives were eluted after (Waters) with a column of µBondasphere C4 (3.9 x 150 mm). concentrated to 2 mL by using Centriprep 10 (Amicon), the and 0.02 % TFA with a flow rate of 0.5 mL/min for 40 min. mutant TPO was further purified by reversed phase HPLC After the TPO fraction (about 30 mL) was approximately 30 min.

biological activity of the derivatives was evaluated by M-07e assay system as previously described. It was found that the all mutants have almost identical activity to that of h6T(1-(63), the reference sample of TPO (See Figures 33 and 34). Centriprep. 10 (Amicon) to approximately 0.1 mg/mL, the For the biological assay, the TPO sample thus phosphate buffer for 2 days. After concentrating with purified was dialyzed two times against 1 L of 10 mM

List of deposits

Escherichia coli (pHT1-231/DH5):

FERM BP-4564 and CCTCC-M95001 Escherichia coli (pEF18S-A2α/DH5): FERM BP-4565 and CCTCC-M95002

FERM BP-4616 and CCTCC-M95003 Escherichia coli (pHGT1/DH5):

FERM BP-4617 AND CCTCC-M95004 Escherichia coli (pHTF1/DH5):

Mouse-Mouse hybridoma P55:

FERM BP-4563 and CCTCC-C95001 CHO28/1/1/3-C6:

FERM BP-4988 and CCTCC-C95004 CHO163T-63-79-C1: FERM BP-4989 and CCTCC-C95005

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SEQUENCE LISTING

(1) GENERAL INFORMATION:

(1) APPLICANT: HIYAZAKI, Birosh:

AKAHORI, Hiromichi KUROKI, Ryota SHINIZU, Toshiyuki MUTO, Takanori (ii) TITLE OF INVENTION: PROTEIN HAVING TPÒ ACTIVITY

(111) NUMBER OF SEQUENCES: 197

(iv) CORMESPONDENCE ADDRESSE:
(A) ADDRESSEE: Marshall, O'Toole, Gerstein, Murxay & Borun
(B) STREET: 6300 Sears Tower, 233 South Macker Drive (A) ADDRESSEE: Marshall, O'Toole, Gere (B) STREET: 6300 Sears Tower, 233 Sout (C) CITY: Chicage (D) STATE: Illinois (E) GOUWIN: United States of America (F) ZIP: 60606-6402

COMPUTER READABLE FORM:

(A) MEDIUM TYPE: Floppy disk (B) COMPUTE: INH PC compatibles (C) OPERATUR SYSTER: PC-DOS/MS-DOS (D) SOFTWARE: Patentin Release #1.0, Version #1.25

CURRENT APPLICATION DATA (41)

(A) APPLICATION NUMBER:
(B) FILING DATE:
(C) CLASSIFICATION:

(vii) PRIOR APPLICATION DATA:

(A) APPLICATION NUMBER: JP 39090/94 (B) FILING DATE: 14-PEB-1994

(vii) PRIOR APPLICATION DATA: (A) APPLICATION NUMBER: JP 79842/94 (B) FILING DATE: 25-MAR-1994

(vii) PRIOR APPLICATION DATA:
(A) APPLICATION NUMBER: JP 155126/94
(B) FILING DATE: 01-JUN-1994

(vii) PRIOR APPLICATION DATA:
(A) APPLICATION NUMBER: JP 167328/94
(B) FILING DATE: 15-JUN-1994

(vii) PRIOR APPLICATION DATA:
(A) APPLICATION NUMBER: JP 227159/94
(B) FILING DATE: 17-AUG-1994

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(A) APPLICATION NUMBER: JP 304167/94
(B) FILING DATE: 01-NOV-1994

(V11)

PRIOR APPLICATION DATA:
(A) APPLICATION NUMBER: JP UNKNOWN
(B) FILING DATE: 28-DEC-1994

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 (B) FILING DATE: 17-AUG-1994

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PRIOR APPLICATION DATA.		
(A) PILING DAIE: 01-DEC-1994 (B) FILING DAIE: 01-DEC-1994	ATION FOR	
PRIOR APPLICATION DATA: (A) APPLICATION NUMBER: JP 193916/94 (B) FILING DATE: 18-AUG-1994	(1) EQUARGE CHARACTER SILES: (B) TYPE: amino acid (D) TOPOLOGY: linear	ica: 1 acids 1
PRIOR APPLICATION DATA: (A) APPLICATION NUMBER: US 08/212,164 (R) PITPE DAFF: 14-MBB-105	(ii) Molecule Type: protein (xi) Sequence Description: Seq ID NO:2:	SEQ ID NO:2:
(y) TALICATION DATA: (A) APPLICATION UNDER: US 08/221,020 (B) FILICATION UNDERS: US 08/221,020 (C) (C) FILICATION UNDERS: US 08/221,020 (C) (C) (C) (C) (C) (C)		Ile Leu Leu Leu Thr -10
(A) APPLICATION UNBER: US 08/278,083 (B) FILING DATE: 20-JUL-1994	Arg Leu Thr Leu Ser Ser Pro Val Pro Pro Ala -5 Leu Aen Lys Leu Leu Arg Aep Ser Tyr Leu Leu 15	Pro Pro Ala Cys Asp Pro Arg Leu 5 10 Tyr Leu Leu His Ser Arg Leu Ser 20 25
PRIOR APPLICATION DATA: (A) APPLICATION NUMBER: US 08/320,300 (B) FILING DATE: 11-007-1994	Asn Pro Leu 35	Pro Val Leu 40
PRIOR APPLICATION DATA: (A) APPLICATION KUMBER: US 08/361,811 (B) FILING DATE: 22-DEC-1994	Val Asp Phe Ser Leu Gly Glu Trp Ly 45 Ala Gln Asp Ile Leu Gly Ala Val Se	Trp Lys Thr Gln Thr Glu Gln Ser Lys 55 Val Ser Leu Leu Leu Glu Gly Val Mer
ATTORNEY/AGENT INFORMATION: (A) NAME: Borun, Michael P. (B) REGISTRANTON NUMBER: 25,447 (C) REFERENCE/DOCKET NUMBER: 01017/13415	Ala Arg Gly Gln Leu Glu Pro	Ser Ser Leu
TELECOMMUNICATION INFORMATION: (A) TELEPRONE: 312/474-6300 (B) TELEPRONE: 312/474-6300	Gly Gln Val Arg Leu 95.	Lou Leu Gly Ala Leu Gln Gly Leu 100 The Tee And And Tee Tee Tee Tee
2) INPORMATION FOR SEQ ID NO:1:	110	oth the lyt Arg Ash lyr Fro Leu Inf 115
SEQUENCE CHARACTERISTICS: (A) LENGTH: 261 base pairs (B) TYPE: nucleic acid	Gln Phe Leu 125 721 THEYDURFFON DON CEA TR MALE.	
(C) STRANDEDNESS: double (D) TOPOLOGY: linear	(1) INFORMATION FOR SEL ID NOIS: (1) SEOUENCE CHARACTERISTICS:	
(ii) molècule type: cona to mena	(£)	pairs
(vi) ORIGINAL SOURCE:(A) ORGANISM: Rattus norvegicus(H) CELL LINE: McA-RH8994	STRANDEDNESS: de TOPOLOGY: lineau LE TYPE: CDNA to	ouble f mRNA
(*i) SEQUENCE DESCRIPTION: SEQ ID NO:1:	ORIGIN	
GTGTACCTG GGTCCTGAAG CCCTTCTTCA CCTGGAIAGA TTCCTTGGCC CACCTGTCCC	(A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: liver	iapiens er
acceacte tetgeagagg tacaaaget caagecetet ceatggeeec aggaaagatt	120 . (xi) SRQUENCE DESCRIPTION: SB	SEQ ID NO:3:
ACCCTGGGCA (CTC GTG GTC ATG CTT CTC CTA ACT Leu Val Val Het Leu Leu Leu Leu Thr	GCA AGG CTA ACG CTG TCC AGC CCG Ala Arg Leu Thr Leu Ser Ser Pro
	GAC CTC CGA	CIC AGT AMA CIG CIT CGT
GA CTA ACT CTG TCC AGC CCA GTT CCT CCC GCC TGT GAC CC FG Leu Thr Lau Ser Ser Pro Val Pro Pro Ala Cys Asp 25		Val Leu Ser Lys Leu Leu Arg Asp 25 30 AGC CAG TGC CCA GAG GTT CAC CAT
	Ser His Val Leu His Ser Arg Leu Se	His Val Leu His Ser Arg Leu Ser Cin Cys Pro Glu Val His Pro

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528 576 580 240 288 336 384 432 480 192 Ala 8 613 160 ព្ព CTG AAC GAG CTC CCA AAC AGG ACT Leu Asn Glu Leu Pro Asn Arg Thr 185 83 F S Xaa Xaa AGC CTC GGA (CTA Val See G GTC NNN Xaa 233 GTC Val 175 65 S Xaa 130 25.5 r 3 Ala 5 th TT By r GTG GAC TIT AGC T A Val Asp Phe Ser L 60 GCA CAG GAC ATT C Ala Gln Asp Ile I 75 GLy CLy GRG Gln 125 ATC 11e ATC Het AGG CGG GCC CCA CCC ACC ACA Arg Arg Ala Pro Pro Thr Thr 170 8 g. GGA ACC C 23 Ala 140 25 ga Ser 1 8 5 4 8 5 8 ŧŝ AAT Phe 155 Ala 8 E3 Arg 5 5 200 ACC AAG C re Cd CCT GCT Pro Ala ATG Het 913 105 105 GAT GTG Val 86c | Ser | 120 Lya GGA AAG CTC ACA Lew Thr CTC Val 23 GAG GAG A Glu Glu T 70 232 95 g CAC His 135 S S ខ្លួ TGC GTC P Cys Val P 165 GGG GCC CTG Gly Ala Leu CTA GTC Leu Val AL AL A Arg 150 553 GAG G1u TCC Ser CAG ATG CCT GTC Pro Val 5 3 Se Ca TG. P P i i 23 ICT Ser 180 5538 X a a 23 TTG CCT ACA C Leu Pro Thr F AAA ACC (£33 CAC HIS ACC Thr 걸 75C Cy8 NNN Xaa EB TTC CAA Phe Gln 1 ខ្លួ Ser th Thr XAA 130 ACA Arg a P TCT G Ser Ser 61y និដ្ឋិនិ Val Val ម្ល 53 KAA

(2) INFORMATION FOR SEQ ID NO:4:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 253 maino acids
(B) TYPE: amino acid
(D) TOPOLOGY: Linear

(ii) MOLECULE TYPE: protein

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:

Gln Cys Pro Glu Val His Pro Leu Pro Thr Pro Val Leu Leu Pro Ala 30 40 Val Asp Phe Ser Leu Gly Glu Trp Lys Thr Gln Met Glu Glu Thr Lys 45 Met Glu Leu Thr Glu Leu Leu Val Val Met Leu Leu Leu Thr Ala -21 -20 Val Leu Ser Lys Leu Leu Arg Asp Ser His Val Leu His Ser Arg Leu Ser 15 Leu Thr Leu Ser Ser Pro Ala Pro Pro Ala Cys Asp Leu Arg P.S.

Ala Cin Asp Ile Leu Gly Ala Val Thr Leu Leu Leu Glu Gly Val Met 60 75 Ala Ala Arg Gly Gin Leu Gly Pro Thr Cys Leu Ser Ser Leu Leu Gly 80 90 Gin Leu Ser Gly Gin Val Arg Leu Leu Ciy Ala Leu Gin Ser Leu Leu Gly Thr Gin Leu Pro Pro Gin Gly Arg Thr Thr Ala His Lys Asp Pro Pro Thr Thr Ala Val Pro Ser Arg Thr Ser Leu Val Leu Thr Leu Asn Phe Thr Pro Asn Ala Ile Phe Leu Ser Phe Gln His Leu Leu Arg Gly Lys Val Arg Phe Leu Met Leu Val Gly Gly Ser Thr Leu Cys Val Arg Arg Ala Ala Ser Ala Arg Thr Thr Gly Ser Gly Leu Leu Lys Trp Gln Gln Gly 190 Phe Arg Ala Lys Ile Pro Gly Leu Leu Asn Gln Thr Ser Arg Ser Leu Asn Glu Leu Pro Asn Arg Thr Ser Gly Leu Leu Glu Thr Asp Gin Ile Pro Gly Tyr Leu Asn Arg Ile His Glu Leu (2) INFORMATION FOR SEQ ID NO:5:

(i) SEQUENCE CHARACTERISTICS:
(A) IENCTH: 576 base pair
(B) TYPE: nuclaic acid
(C) SYRANDENESS: double
(D) TOPOLOGY: linear (11) MOLECULE TYPE: CDNA to MRNA

(A) ORGANISH: Homo Sapiens (F) TISSUE TYPE: liver (vi) ORIGINAL SOURCE:

(x1) SEQUENCE DESCRIPTION: SEQ ID NO:5:

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CAA ACC TCC AGG Gln Thr Ser Arg TTC 9. S. S. GAA CTC 1 Glu Leu 1 30 e 3 AC of the 1 CAC His CGC AGG CTG AAC C Leu Asn C CTG AAC AGG A Leu Asn Arg I 25 TCA GAC ACA G Ser CCT GGA CCC 7 reg Cig AGA GCC AAG ATT CCT GGT Arg Ala Lys Ile Pro Gly ICC CTG GAC CAA ATC CCC GGA TAC Ser Leu Asp Gln Ile Pro Gly Tyr 20 GGA CTC TTT G GL GL Ser 3 TCC ACT CGT G Thr Arg G ATT 11e GGA TTC 2 £ 5 GCC CCC AST ğ

191 239 អ្ន ដ 8 3 CCA Pro Th. i Fer £ 5 £ £ Ser 61y S CAT The Popular Pro Pro Ser Aca Thr ដូដូ Ser id H. g g P S 5 gg 20 950 Ash Leu

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	287	335	383	432	492	552	576		
75 75	AGG CTC TTC CCT CCT AGC TTG CCC AGC CCT GTG GTC Thr Leu Phe Pro Leu Pro Thr Pro Val Val 95	CAC CCC CTG CTT CCT GAC CCT TCT GCT CCA AGG CCC ACC CCT His Pro Leu Leu Pro Asp Pro Ser Ala Pro Thr Pro Thr Pro 110	CT CIT CIA AAC ACA ICC IAC ACC CAC ICC CAG AAI CIG ICT Pro Leu Leu Aen Ihr Ser Tyr Ihr Hie Ser Gin Aen Leu Ser 126	56 IAAGGITCIC AGACACIGCG GACAICAGCA IIGICICAIG 1y 30	TACAGCTCCC TTCCCTGCAG GGGGCCCCTG GGAGACAACT GGACAAGAIT TCCTACTTTC	TCCTGABACC CARAGCCCTG GTARARGGGA TACACAGGAC TGARARGGGA ATCALTITTC	ACTGIACAIT AIAAACCIIC AGAA	(2) INFORMATION FOR SEQ ID NO:6:	(i) SEQUENCE CHARACTERISTICS:
65	CAG TAT A Gln Tyr T 80	CAG CTC C Gln Leu H	ACC AGC CCT Thr Ser Pro	CAG GAA GGG 7 GIn Glu Gly 130	TACAGCTCC	TCCTGAAAC	ACTGTACAI	(2) INPOF	i)

Ala Ala Arg Cly Cln Leu Gly Pro Thr Cys Leu Ser Leu Leu Gly 80 80 90 Leu Gly Thr Gln Leu Pro Pro Gln Gly Arg Thr Thr Ala His Lys Asp 110 Phe Leu Met Leu Val Gly Gly Ser Thr Leu Cys Val Arg Arg Ala 155 Gin Cys Pro Glu Val His Pro Leu Pro Thr Pro Val Leu Leu Pro Ala 36 40 Ala Gln Asp Ila Leu Gly Ala Val Thr Leu Leu Glu Gly Val Met 60 75 Pro Aan Ala Ile Phe Leu Ser Phe Gln His Leu Leu Arg Gly Lys Val 125 Het Glu Leu Thr Glu Leu Leu Val Val Het Leu Leu Leu Thr Ala -21 -20 Leu Ser Lys Leu Leu Arg Asp Ser His Val Leu His Ser Arg Leu Ser 15 Val Asp Phe Ser Leu Gly Glu Trp Lys Thr Gln Het Glu Glu Thr Lys 45 Gin Leu Ser Cly Gin Val Arg Leu Leu Ceu Gly Ala Leu Gin Ser Leu 100 Arg Leu Thr Leu Ser Ser Pro Ala Pro Pro Ala Cys Asp Leu Arg Val -5 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:6: (ii) MOLECULE TYPE: protein Arg 140

Asp lie ser Ser Gly Thr Ser Asp Thr Gly Ser Leu Pro Pro Asn Leu 260 Gln Pro Gly Tyr Ser Pro Ser Pro Thr His Pro Pro Thr Gly Gln Tyr 270 Leu Phe Pro Leu Pro Pro Thr Leu Pro Thr Pro Val Val Gln Leu 285 His Pro Leu Leu Pro Asp Pro Ser Ala Pro Thr Pro Thr Pro Thr Ser 310 Pro Leu Leu Asn Thr Ser Tyr Thr His Ser Gln Asn Leu Ser Gln Glu 320 Ala Ser Ala Arg Thr Thr Gly Ser Gly Leu Leu Lys Trp Gln Gly Ala 190 Asp Gln Ile Pro Gly Tyr Leu Asn Arg Ile His Glu Leu Leu Asn Gly 220 Thr Arg Gly Leu Phe Pro Gly Pro Ser Arg Arg Thr Leu Gly Ala Pro 240 250 Aen Glu Leu Pro Aen Arg Thr Ser Gly Leu Leu Glu Thr Aen Phe Thr 185 Leu Phe Arg Ala Lys Ile Pro Gly Leu Leu Asn Gln Thr Ser Arg Ser 216 216 Pro Pro Thr Thr Ala Val Pro Ser Arg Thr Ser Leu Val Leu Thr 160 (i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 1721 base pairs
(B) TYPE: nucleic acid
(C) STRANDENESS: double
(D) TOPOLOGY: linear (vi) ORIGINAL SOURCE:
(A) ORGANISH: Homo Sapiens
(F) IISSUE TYPE: liver (ii) HOLECULE TYPS: CONA to mRNA (2) INFORMATION FOR SEQ ID NO:7: IP. Ç,

GOGGCACGAG GGGGTGTCT GGCTGGCGTG GCTCCTGTT TGGGGCCTCT CCCCTGAATC GAM ING CIC CIC GIG GIC AIG CIT CIC CIA ACT GCA AGG CIA AGG GIU Leu Leu Leu Val Val Het Leu Leu Leu Thr Ala Arg Leu Thr Leu -15 -15 15 gg GAG G1.u Thr Thr AAA Lys 1 TGC CCA C CTICCISGGG CCATGGAGGC CAGACAGACA CCCCGGCCAG A ATG GAG CTG
Het Glu Leu
-21 -20 AGT CTG AGC CAG 1 Leu Ser Gln G 53 GTC 8 5 S (xi) SEQUENCE DESCRIPTION: SEQ ID NO:7: 253 CAC AGC AGA TGT GAC (Cys Asp) EŠ k g CCT CCT G Pro Pro A TCC CAT GTC Ser His Val 1 A La 2000 Asp Ser Ser Arg Arg 53

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1350 1410 1470

1590 1650 1710

20 25 30		Leu Pro Pro Thr Leu Pro Thr Pro Val Val Gln Leu His Pro Leu Leu
71T CAC CCT TTG CCT ACA CCT GTC CTG CTG CCT GCT GTG GAC TTT AGC /al His Pro Leu Pro Thr Pro Val Leu Leu Pro Ala Val Asp Phe Ser 40	305	TCT GCT CCA AGG CCC AGC CCT AGG AGG Ser Ala Pro Thr Pro Thr Ser
ITG GGA GAA TGG AAA ACC CAG ATG GAG GAG ACC AAG GCA CAG GAC AIT .au gly glu Tip Lyb Thi Gln Het glu glu Thi Lyb Ala gln Abp Ile 55	353	305 TCC TAC ACC CAC TCC CAG AAI CTG TCT CAG GAA GGG TAAN Ser Tyr Thr His Ser Gln Aan Leu Ser Gln Glu Gly
TIG GGA GCA GTG ACC CTI CTG CTG GAG GGA GTG ATG GCA GGA CGG GGA GOU GJY Ala Val The Lou Lou Lou Glu Gly Val Hot Ala Ala Arg Gly 65	401	325 ACAICAGCA ITGICICGIG IACAGCI
TAA CTG GGA CCC ACT TGG CTG TCG CTG GGG CAG CTT TCT GGA Lau Lau Gly Pro Thr Cys Lau Ser Ser Leu Lau Gly Gln Leu Ser Gly 80 90 95	449	GGRGRCRACT GGRCRAGATT TCCTRCTTTC TCCTGARACC CRARGCCCTG GTALARGGGA TACATTTTC ACTGIACATT ATRARCCTTC AGRAGCTATT
AG GTC CGT CTC CTT GGG GCC CTG CAG AGC CTC CTT GGA ACC CAG iin Val Arg Leu Leu Leu Gly Ala Leu Gln Ser Leu Leu Gly Thr Gln 105	497	ITITAAGCI AICAGCAATA CTCATCAGAG CAGCTAGGTC ITTGGTCTAT ITTGTGCAGA AAITIGGAAC TCACTGAITC TCIACAIGCI CITITICIGI GAIAACICIG CAAAGGCCTG
TIT CCT CCA CAG GGC AGG ACC ACA GGT CAC AAG GAT CCC AAT GCC ATC Leu Pro Pro Gln Gly Arg Thr Thr Ala His Lys Asp Pro Asn Ala Ile 115	545	GECTEGECCIG GENGTICANE AGAGGGAGAG AETAACETIG AGTCAGAAAA CAGAGAAAGG GTAATTICCI TIGCTICAAA ITCAAGGCCI ICCAAGGCCC CCAICCCCII IACTAICAIT
TIC CTG AGC TIC CAA GAC CTG CTC CGA GGA AAG GTG CGI TIC CTG ATG Phe Leu Ser Phe Gin His Leu Leu Arg Gly Lys Val Arg Phe Leu Met 130	593	CICACTOGGA CICTGAICCC ATAITCITAA CAGAICTITA CICTIGAGAA ATGAATAAGC TTTCICTCAG AAATGCTGTC CCTATACACT AGACAAAACT GAAAAAAAA AAAAAAAAA
ITT GIA GGA GGG TCC ACC GTC AGG GGG GCG CCA CCC ACA AGA VAl Gly Gly Ser Thr Leu Cys Val Arg Arg Ala Pro Pro Thr Thr 145	641	AMAMAMAM A (2) Information for SEQ ID No:8:
ICT GTC CCC AGC AGC TCT CTA GTC CTC ACA CTG AAC GAG CTC CCA Na Val Pro Ser Arg Thr Ser Leu Val Leu Thr Leu Aen Glu Leu Pro 160 170 170	689	(i) SEQUENCE CHARACTERISTICS: (a) LENGTH: 4506 base pairs (b) TYPE: nucleic acid (c) STRANDEDNESS: double
NAC AGG ACT TCT GGA TTG TTG GAG ACA AAC TTC ACT GCC TCA GCC AGA NAN ARG Thr Ser Gly Leu Leu Glu Thr Asn Phe Thr Ala Ser Ala Arg 180	737	(D) TOPOLOGI: Linear (ii) MOLECULE TYPE: Genomic DNA
NCA ACT GGC TCT GGG CTT CTG AAG TGG CAG GGA TTC AGA GCC AAG Thr Thr Gly Ser Gly Leu Leu Lys Trp Gln Gln Gly Phe Arg Ala Lys 195	785	(vi) ORIGINAL SOURCE: (A) ORGANISH: Homo Saplens (G) CELL TYPE: peripheral leukocytes
NIT OCT GGT CIG CIG AAC CAA ACC TCC AGG ICC CIG GAC CAA AIC CCC lle Pro Gly Leu Leu Asn Gln Thr Ser Arg Ser Leu Asp Gln Ile Pro 210	833	(*1) SEQUENCE DESCRIPTION: SEQ ID NO:8: GAATTCAGGG CITTGGGAGT TCCAGGCTGG TCAGGATCTC AAGCCCTGCC CAGGATCTGT
GGA TAC CTG AAC AGG ATA CAC GAA CTC TTG AAT GGA ACT CGT GGA CTC 31y Tyr Leu Abn Arg Ile His Glu Leu Leu Abn Gly Thr Arg Gly Leu 235	881	TCACCCTGCC AGGCAGTCTC TICCTAGAAA CTTGGTTAAA TGTTCACTT TCTTGCTACT ITCAGGAIAG AITCCTCACC CTTGGCCCGC CTTTGCCCCA CCCTACTCTG CCCAGAAGTG
ITI CCI GGA CCC ICA CGC AGG ACC CTA GGA GCC CCG GAC AII ICC ICA Phe Pro Gly Pro Ser Arg Arg Thr Leu Gly Ala Pro Asp lle Ser Ser 240 250 245 245 245 250 250 250 250 250 250 250 250 250 25		CAAGAGCTIA AGCGCCTCC ATGGCCCCAG GAAGATTCA GGGGAGAGGC CCCAAACAGG GAGCCAGGCC AGCCAGACAC CCCGGCCAGA ATG GAG CTG ACT G GTGAGAACAC Mgt Glu Leu Thr
56A ACA TCA GAC ACA GGC TCC CTA CCC AAC CTC CAG CCT GGA TAT 51y Thr Ser Asp Thr Gly Ser Leu Pro Pro Asn Leu Gln Pro Gly Tyr 265	716	-21 -20 Acctergegg Ctrgggcecelt atggradert gacagraggg gagagarang ggagacacgc
TOT CCT TCC CCA ACC CAT CCT CCT ACT GGA CAG TAT ACG CTC TTC CCT Ser Pro Thr His Pro Pro Thr Gly Gln Tyr Thr Leu Phe Pro 280	1025	TUCAGGGGG AGGAGCTGG GGGAACCCAT TCTCCCAAAA AIAAGGGGTC TGAGGGGTGG ATTCCCTGGG TTTCAGGTCT GGGTCCTGAA TGGGAATTCC TGGAATACCA GCTGACAATG
כדד ככא כככ אככ דדם כככ אככ ככד קדם פדכ כאה כדכ כאכ כככ כדם כדד	1073	ATTICCICCT CAICTITCAA CCICACTCT CCICAICIAA G AA IIG CTC CTC GIu Leu Leu Leu

120 180 240 293

2145 2205 2265 2325 2385 2445 2505 2565 2625

CAATAGITIA AAAAACIAAA

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NICIATCCIC AAGAACCCIA GCGICCCIIC IICCIICAGG ACIGAGICAG GGAAGAAGG GGGATICICC IGCTCAGT ICCCAAGIAG CIGGGAIIÀC AGGIGCCCAC GGCCAIGCCC AGCIAAITIT IGIAITITIG GIAGAGAIGG GGITICACCA IGIIGGCCAG GCIGAICITG AACTCCTGAC CTCAGGTGAI CCACCTGCCI CAGCCTCCCA AAGTGCTGGG AITACAGGGG TEAGCCACTG CACCCAGCCT TCATTCAGTT TAAAAATCAA ATGATCCTAA GGTTTTGCAG cagcaacgta agaaaaagg agctettete actgaaacca agtgtaagae caggetggae GAATICCIGC CCIGGGIGGG ACCTIGGICC IGICCAGIIC ICAGCCIGIA IGAIICACIC ccaggotigga grocagiggo algaitteaa citeacidaa cotcagcoro coggaiicaa cagaaagagt aaaittgcag cactagaacc aagaggtaaa agctgtaaca gggcagaitt IAGAGGACAC GGGAGITITI GAAGCAGAGG CTGATGACCA GCTGTGGGGA GACTGTGAAG IGCIGGCIAC ICCIAAGGCI CCCCACCCCC IIIIAGIGIG CCCIIIGAGG CAGIGCGCII GTAAGTCCCC AGTCAAGGA TCTGTAGAAA CTGTTCTTTT CTGACTCAGT CCCCCTAGAA GACCTGAGGG AAGAAGGGCT CTICCAGGGA GCTCAAGGGC AGAAGAGCTG ATCTACTAAG CICICITICCA TCICITICIC AG GAG GAG AAG GCA CAG GAC AIT CIG GGA Glu Glu Thr Lys Ala Gln Asp Ils Leu Gly 60 60 65 AGTGCTCCCT GCCAGCCACA ATGCCTGGGT ACTGGCATCC TGTCTTTCCT ACTTAGACAA GGGAGGCCTG AGAICTGGCC CIGGIGITIG GCCTCAGGAC CAICCICIGC CCICAG GCT CAC AAG GAT CCC AAT ALS His Lys Asp Pro Ash 120 GAG GGA GTG ATG GCA GCA CGG Glu Gly Val Met Ala Ala Arg AAG GTG Lys Val SCAGCCTGAA CAGAAAGAGA CTAGAAGCAT GTTTTATGGG CICAIACCIA CAIIIAGIII AITIAIIAII AIIAIIIGAG TITICAICCE 550 E3 Arg 현취당 61,4 61,4 re Cig g g AGG ž 3 2538 AGC Ser A 26 Vel Gr GTC Val CAGITCCIAI GGGTCCCTIC IAGICCIIIC 25.0 10.0 10.0 53 CAG GGC AGG ACC ACA Gln Gly Arg Thr Thr 115 CTC 135 TGC £ 3 Ser 233 55.3 150 CH Ser 19 CTC CTC feu Leu 70 CTC TCA Leu Ser GGG GCC Gly Ala CAC Acc Thr 165 7 tr 35 JCC Ser AGA ACC CTT Thr Leu 76C Cys E3 THE BHG 99.5 AGC F F33 CCT CCA AGC Ser 130 8 2 ន្តិដ GCA GTG A 200 OGT CTC CTC Len CTC GTA Val G G E3 TTC Phe E 3 Ala 160 672 2025 2085 573 621 792 852 912 1245 1305 1485 1545 1665 1785 1845 1905 1965 732 961 6001 1065 1125 1185 1365 1425 605 725 GCTTGGCCAC CCTAACCCAA TCTACATTCA CCTATGATGA TAGCCTGTGG ATAAGATGAT GITIGATGIT IAGCAICCCC AITGIGGAAA IGCICGIACA JAAGACHIAT GCTAATITAI TAAGAGGGAC CATAITAAAC TAACATGIGI CTAGAAAGCA CTAITCTICC CAINITIGICC CCACCIACTG AICACACTCI CTGACAAGGA TTATICITCA CHAIRCAGCC CSCAITIAAA AGCICICGIC IAGAGAIAGI ACICAIGGAG CITAITAGGC TACCATAGCT CTCTCTATTT CAGCTCCCTT CTCCCCCAC GGCTTGCAGG TCCAATATGT GAATAGATTT GAAGCTGAAC ACCATGAAAA GCTGGAGAGA aagcaagact catatgecat ccacagatga cacaaagctg ggaagtacca ctaaaataac aaaagactga atcaagattc aaatcactga aagactaggt caaaaacaag gtgaaacaac AGAGATATAA ACTICIACAI GIGGGCGGG GGCTCACGCC IGTAAICCCA GCACTITGGG NGCCGAGGC AGGCAGATCA CCTGAGGGCA GGAGTTTGAG AGCAGCCTGG CCAACATGGC GANACCCCGT CTCTACTANG ANTACAANT TAGCCCCGCA TGGTAGTGCA TGCCTGTAAT GGAGGTTGTA aaagaaaa aaaattetac atgtotaaat taatgagtaa agtoctatte cagettega ACGARARGGA ICTGAGAGAA ITAAAITGCC CCCAAACTTA CCAIGTAACA TTACTGAAGC TGCIAITCTT COCTAACTG GIAAGACACC CATACTCCCA GGAAGACACC ATCACTTCCT CTAACTCCTT CTCACCTTCA FIGAGCTGAG ATCATGCCAA TGCACTCCAG CCTGGGTGAC AAGAGCAAAA CTCGGTCTCA CCT GCT GTG GAC TIT AGC TIG GGA GAA TGG AAA ACC Pro Ala Val Asp Phe Ser Leu Gly Glu Trp Lys Thr 45 ACG CTG TCC AGC CCG GCT Thr Leu Ser Ser Pro Ala CTC AGT AAA CTG CTT CGT GAC TCC Leu Ser Lys Leu Leu Arg Asp Ser CAG ATG GTAAGAAAGC CATCCCTAAC CTTGGCTTCC CTAAGICCTG ICTTCAGITT Gla Met 55 CAATCITITI CAACAG ACC CAG TGC CCA GAG GII CAC CCI I'G CCI ACA Ser Gln Cys Pro Glu Val His Pro Leu Pro Ihr 30 CTG GTGAGAACTC CCAACATTAT CCCCTTTATC Leu GAACTCTAIT CCGAGIGGAC TACACTIAAA TAIACTGGCC IGAACACCGG TTAAAAATAT AATCGCTCAI GGCCAIGCCI TTGACCTAIT CCIGIICAGI CIICIIAAAI CCCAGCTACT TGGAAGGCTG AAGCAGGAGA ATCCCTTGAA CCCAGGAGGT SCCACAAIGC CCIGCTICCA ICATTIAAGC CICIGGCCCI AGCACTICCI CCCACTGCTI CCCATGGAIT CTCCAACAIT CITGAGCTIT 63 Thr Ale Arg I TGT GAC CTC CGA GTC Cys Asp Leu Arg Val 10 E 3 AGA Arg

CTG CTG Leu Leu

CCT GTC Pro Val

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GGA CAA C

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TCT GGA CAG G

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3235 3295 3351 3399 3447

ATC

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GCC Ala 155

AAAGCIAGIA AITCITGICI

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CCA Pro 175

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GAG Clu 1

Asn

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cac Gla

GGA ACC Gly Thr G

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	AND THE SET CLY LOW LOW CHE THE MEN PIN THE ALE SET ALE AND THE ALE SET ALE AND THE ALE THE GLASS OF CAG
ACT GGC TCT GGG CTT CTG AAG TGG CAG GGA TTC AGA Thr Gly Ser Gly Leu Leu Lys Trp Gln Gln Gly Phe Agg Thr Gly Ser Gly Leu Leu Lys Trp Gln Gln Gly Phe Agg CCT GGT CTC CTG AAC CAA ACC TCC AGG TCC CTG GAC CAA Pro Gly Leu Leu Asn Gln Thr Ser Arg Ser Leu App Gln TAC CTG AAC AGG ATA CAA ACC TCC AGG TCC CTG GAC CAA TYC CTG AAC AGG ATA CAC GAA CTC TTG AAT GGA ACT CCT Tyr Leu Asn Arg lle His Glu Leu Leu Asn Gly Thr Arg 215 ACA TCA GAC AGG ATA CTC CTA GGA CCC CG GAC ATT Pro Gly Pro Ser Arg Arg Thr Leu Gly Ala Pro Asp Ile 245 ACA TCA GAC AGG CTC CTC CCA CCC CAC CTC CG TAC GAC AGG CTC CTC CCA CCC CAC CTC CAC Thr Ser Asp Thr Gly Ser Leu Pro Pro Asn Leu Gln Pro CTY CC CAC ACC CTC CTC CCA CCC CAC CTC CAC TCC CCC ACC CTC CTC CCA CCC CAC CTC CAC TCC CCA CCC TCT CCC ACC CCT CAC CCC CTC CT	ACT CGC TOT GGG CTT TOTA AG TGG CAG GGA TTC AGA Thr Gly Ser Gly Leu Leu Lys Typ Gln Gln Gly Phe Aga Thr Gly Ser Gly Leu Leu Lys Typ Gln Gln Gly Phe Aga Thr Gly Ser Gly Leu Leu Ann Gln Thr Ser Arg Ser Leu App Gln Tac CTG AAC ACC CAA ACC CTC AGG CC CTG GAC CAT Typ Leu Ann Arg lle His Glu Leu Leu Ann Gly Thr Arg Typ Leu Ann Arg lle His Glu Leu Leu Ann Gly Thr Arg Typ Leu Ann Arg lle His Glu Leu Leu Ann Gly Thr Arg Typ Cly Pro Ser Arg Arg Thr Leu Gly Ala Pro App lle Thr Ser Ang Thr His Glu Leu CC CC GAC ATT Pro Gly Pro Ser Arg Arg Thr Leu Gly Ala Pro App lle Thr Ser App Thr Gly Ser Leu Pro Pro Ann Leu Gln Pro CCT GCA CC CAC CC CT CC CC CC CC CC Thr Ser App Thr Gly Ser Leu Pro Thr Gly Gln Tyr Thr Leu 245 ACA TCA ACC CAC CCT CCT CCA CC CC CC CC Typ Pro Ser Ala Pro Thr Pro Thr Gly Gln Tyr Thr Leu 250 CCT CC CCA ACC CT CC CC CC CC ACC CC CC Typ Pro Ser Ala Pro Thr Pro Thr Gly Gln Leu His Pro 250 GAC CCT TCT GCT CC CAC CCT ACT GCA CCT CCT Asp Pro Ser Ala Pro Thr Pro Thr Pro Thr Ser Pro Leu 305 AND Thr His Ser Gln Ash CCT CCT CAC CAC CCT TTC TAC ACC CAC TCT CCT CAC CAC CTT TTC TAC ACC CAC TCT CCT CAC CAC CTT TTC TAC ACC CAC TCT CTTACATATT TCTACATACTT TTGCAAC TGACAGATT TCTACATACT TCCTGAAAAC TTGCACC GACATACACT TCTACATACT TTTTTTTTT TATACTT ATAAACCTT TTGCAAC TGACAGATT TCTACATACT TCCTGAAAAC TTGCACC GACATACATAC TCTACATACT TTTTTTTTTT
AND	

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 416 base pair
(B) TIPE: nucleic acid
(C) STRANNEDNESS: double
(D) TOPOLOGY: linear

(!!) MOLECULE TYPE: synthetic DNA

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ID NO:9:
SEO
DESCRIPTION:
SEQUENCE
(x r)

CTAGAAAAA	CCAAGGAGGT	CTAGAAAAAA CCAAGGAGGI AATAAATAAI GAGCCCGGGT CCGCCAGCTT GTGACCTTCG	GAGCCCGGCT	CCCCAGCTT	GTGACCTTCG	9
TGTTCTGTCT	AAACTGCTTC	TETICIGICI AAACIGCTIC GCGACICICA CGIGCTGCAC ICICGICIGI CCCAGIGCCC	corecrecae	тстсетстет	CCCAGTGCCC	120
GGAAGTTCAC	COCCTGCCGA	GGAAGTTCAC CCGCTGCCGA CCCCGGTTCT GCTTCCGGCT GTCGACTTCT CCCTGGGTGA	GCTTCCGGCT	GTCGACTTCT	CCCTGGGTGA	180
ATGGAAAACC	CAGATGGAAG	ATGGAAAACC CAGATGGAAG AGACCAAACÇ TCAGGACATC CTGGGTGCAG TAACTCTGCT	TCAGGACATC	CTGGGTGCAG	TAACTCTGCT	240
TCTGGAAGGC	GTTATGGCTG	ICTGGAAGGC GITAIGGCIG CACGIGGCCA GCIIGGCCCG ACCIGCCIGI CIICCCIGCI	GCTTGGCCCG	ACCTGCCTGT	CTTCCCTGCT	300
TGGCCAGCTG	TCTGGCCAGG	rescensere rerescense rresrerer seresseer erschstere rserrssene	GCTCGGCCCT	CTCCAGTCTC	TGCTTGGCAC	360
ccaecreece	CCACAGGGCC	CCAGCTGCCG CCACAGGGCC GIACCACIGC ICACAAGGAI CCGAACGCIA ICTICC	TCACAAGGAT	CCGAACGCTA	TOTTOG	416
(2) INFORMA	TION FOR SE	(2) INFORMATION FOR SEQ ID NO:10:				
(1)	SEQUENCE CE (A) LENGI (B) TYPE: (D) TOPOL	(1) SEQUENCE CHARACTERISTICS: (A) LENGTH: 179 amino acids (B) TYPE: amino acid (D) TOPOLOGY: linear	CS: o acids			
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(xi) SEQUENCE DESCRIPTION: SEQ ID NO:10: (ii) MOLECULE TYPE: protein

Gin Cya Pro Giu Val His Pro Leu Pro Thr Pro Val Leu Leu Pro Ala Val Amp Phe Ser Leu Gly Glu Trp Lym Thr Gln Met Glu Glu Thr Lym 45 55 Ala Gin Asp ile Leu Gly Ala Val Thr Leu Leu Leu Glu Gly Val Met 60 75 Leu Gly Thr Gin Leu Pro Pro Gin Gly Arg Thr Thr Ala His Lys Asp Arg Phe Leu Met Leu Val Gly Gly Ser Thr Leu Cys Val Arg Arg Ala Pro Pro Thr Thr Ala Val Pro Ser Arg Thr Ser Leu Val Leu Thr Leu Leu Val Pro Arg Gly Ser Pro Ala Pro Pro Ala Cys Asp Leu Arg Val -5 Leu Ser Lys Leu Leu Arg Asp Ser His Val Leu His Ser Arg Leu Ser Ala Ala Arg Gly Gln Leu Gly Pro Thr Cys Leu Ser Ser Leu Leu Gly 80 80 Gin Leu Ser Gly Gin Val Arg Leu Leu Leu Gly Ala Leu Gin Ser Leu Pro Asn Ala Ile Phe Leu Ser Phe Gln His Leu Leu Arg Gly Lys Val

Asn Glu Leu

(2) INFORMATION FOR SEQ ID NO:11:

(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 535 base pairs (B) TYPE: nucleic acid

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			52	100	148	196	244	292	340	388	436	484	533	535	
	(*1) ORIGINAL SOURCE: (A) ORGANISH: Homo Sapiens	(*1) SEQUENCE DESCRIPTION: SEQ ID NO:11:	CTRGRARANA CCARGGRGCT ANTRANTA ATG ANA GCA CCT GTA CCA CCT GCA Met Lys ala Pro Val Pro Pro Ala -2 1	TGT GAT TTA GGG GTC CTG TCT AAA CTG CTG CGC GAC TCT CAC GTG CTG Cys Asp Leu Arg Val Leu Ser Lys Leu Leu Arg Asp Ser His Val Leu 10 11 15 12 12 12 12 13 15 15 15 15 15 15 15 15 15 15 15 15 15	CAC TCT CGT CTC TCC CAG TGC CCG GAA GTT CAC CCG CTG CCG ACC CCG His Ser Arg Leu Ser Gln Cys Pro Glu Val His Pro Lau Pro Thr Pro 35	GTI CTG CTI CCG GCT GTC GAC TTC TCC CTG GGT GAA TGG AAA ACC CAG Val Leu Leu Pro Ala Val Asp Phe Ser Leu Gly Glu Trp Lys Thr Gln 40 50	ATG GAA GAG ACC AAA GCT CAG GAC ATC CTG GGT GCA GTA ACT CTG CTT Het Glu Glu Thr Lys Ala Gln Asp Ile Leu Gly Ala Val Thr Leu Leu SS 65	CTG GAA GGC GTT AIG GCT GCA CGT GGC CAG CTT GGC CGG ACC IGC CTG Leu Glu Gly Val Met Ala Ala Arg Gly Gln Leu Gly Pro Thr Cys Leu 75	ICT ICC CIG CIT GGC CAG CIG ICT GGC CAG GII CGI CIG CIG CIG GGC Ser Ser Leu Leu Gly Gln Leu Ser Gly Gln Val Arg Leu Leu Leu Gly 95	GCT CTG CAG TCT GTG CTT GGC ACC CAG CTG CCG CAG GGC CGT ACC Ala Leu Gln Ser Leu Leu Gly Thr Gln Leu Pro Pro Gln Gly Arg Thr 105	ACT GCT CAC AAG GAT CCG AAC GCT ATC TTC CTG TCT TTC CAG CAC CTG Thr Ala His Lys Asp Pro Asn Ala Ila Phe Leu Ser Phe Gln His Leu 120	CTG CGT GGC AAA GTT CGT TTC CTG ATG CTG GGT GGC GGT TCT ACC CTG Leu Arg Gly Lys Val Arg Phe Leu Met Leu Val Gly Gly Ser Thr Leu 135	IGC GTT CGT CGG GCG CCG CCA ACC ACT GCT GTT CCG TCT TAATGAAAGC Cys Val Arg Arg Ala Pro Pro Thr The Val Pro Ser 155	E	(2) INFORMATION FOR SEQ ID NO:12:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 515 base pairs
(B) TYPE: Nucleic acid
(C) STRANDEDNESS: double
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: synthetic DNA

(vi) ORIGINAL SOURCE:

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Sapiene	
Homo	
ORGANISM:	
-	

NO: 12:
B
SEQ
DESCRIPTION:
SEQUENCE
(XL)

52	100	148	196	. 244	292	340	388	436	484	533	535
CTAGAAAAA CCAAGGAGGT AATAAATA ATG AAA TCT CCT GCA CCT GCA Het Lys Ser Pro als Pro Als -2 1	GAT TIA GGG GTC CTG TCT AAA CTG CTG GGC GAC TCT CAC GTG CTG ASP Leu Arg Val Leu Ser Lys Leu Leu Arg Asp Ser His Val Leu 10 10 15 15 15 15 15 15 15 15 15 15 15 15 15	TOT COT CTG TCC CAG TCC CCG GAA GTT CAC CCG CTG CCG ACC CCG 146 Ser Arg Leu Ser Gin Cys Pro Giu Val His Pro Leu Pro Thr Pro 25	CTG CTT CCG GCT GTC GAC ITC TCC CTG GGT GAA TGG AAA ACC CAG 196 Leu Leu Pro Ala Val Asp Phe Ser Leu Gly Glu Trp Lyg Thr Gln 40	GAA GAG ACC AAA GCT CAG GAC ATC CTG GCT GCA GTA ACT CTG CTT 24. Glu Glu Thr Lys Ala Gln Asp Ile Leu Gly Ala Val Thr Leu Leu 60 65	GAA GGC GTT ATG GCT GCC CAG CTT GGC CCG ACC TGC CTG Glu Gly Val Met Ala Ala Arg Gly Gln Leu Gly Pro Thr Cys Leu 75	ICC CTG CTI GGC CAG CTG TCC CGC CTG CTG CTG CTG CGC 340 Ser Leu Leu Gly Gln Leu Ser Gly Gln Val Arg Leu Leu Leu Gly 90 95 100	CTG CAG TCT CTG CTT GGC ACC CAG CTG CCG CCG ACC 388 Leu Gln Ser Leu Leu Gly Thr Gln Leu Pro Pro Gln Gly Arg Thr 105	GGT CAC AAG GAT CGG AAC GCT ATC TTC CTG TCT TTC CAG CAC CTG 43 Ala His Lys Asp Pro Asn Ala Ile Phe Leu Ser Phe Gln His Leu 120	CGT GGC AAA GTT CGT TTC CTG ATG CTG GGT GGC GGT TCT ACC CTG 48. Arg Gly Lys Val Arg Phe Leu Het Leu Val Gly Gly Ser Thr Leu 140	GIT CGT CGG CGG CCG CCA ACC ACT GCT CTT CCG TCT TAATGAAAGC Val Arg Arg Ala Pro Pro Thr Thr Ala Val Pro Ser 155	83
CTAGN	TGT G	CAC TO Bis So	12 y	ATG G	CTG Lea G	Ser S	GCT C	ACT G Thr A	CTG O Leu A 135	TGC G Cys V	£

(2) INFORMATION FOR SEQ ID NO:13:

(1) SEQUENCE CHARACTERISTICS:
(A) LEWGTH: 555 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: double
(D) TOPOLOGY: linear

(11) MOLECULE TYPE: CDNA to MRNA

(vi) ORIGINAL SOURCE:
(A) ORGANISM: Homo Sapiens
(F) IISSUE TYPE: liver

(*1) SEQUENCE DESCRIPTION: SEQ ID NO:13:

ATG GAG CTG ACT GAA TTG CTC CTC GTG GTC ATG CTT CTC CTA ACT GCA

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95 E 3 950 17 A th Leu ន្តិន 535 re 3 82 CAT His 230 A GC re d 150 33 TCT Cic Ci AGC 5 5 75. Cys 85 P P CTC Leu CTC Leu Arg P P Se d ABC ABD ATC 11e Val 참 CAC ACC Thr 165 E CE CGT 245 950 CAT Hi.B 8 5 PA Lys Acc Thr Arg 밝 55.9 95°C CAG Gln TCT 180 180 180 66c 61y Arg AGC 25 th 25 Thr. Ser 35.53 55 GTA Val ng g re d 95 115 115 Phe 69.4 61.4 TCT ACC AGC Ser 195 CTC Leu AAC 500 GAT Pro 275 Leu Leu Arg Asp ខ្លួន 2538 Ala Ala Arg S C O 130 130 900 61y Pro or ACC Thr 950 C17 210 210 95.5 AGC 5 3 AGC 95.5 GTT Val 145 2 CC E CAC St. 617 658 t a Va TT re 3 GTT Val Arg CGT ACC Thr 97 C TAT Tyr 225 55 5 뀱 7 CG Ala 1 MAC ATC 11e 5 3 2 8528 re d 7 K 660 614 61.y Ser Val re Ci 5 th TTC Phe E G Phe 240 g cac MTC 275 175 175 SP. Ser ATC Ile 55.5 61y 95 A St P Act Arg A Eys Eys 8 5 E 3 TAT. TCT Ser 255 CAG GAC Gln Asp CCA ACC 15 g 8 K ATC 11e F 63 Ser 111 Ala 9 K 60 ន្លដ្ឋន F ag ខ្លួ Giy Giy AGC 213 270 270 Ser Lys AAC ABB GAG Ser 25 cg Glo Gla 225 GAC ABP A Sca E 3 61.7 61.7 TTC Arg ATC ន្តដ ren 95 P 48 A La GAG Gla E 3 5 5 Arg 140 3 5 ABD A GCG 뀱 GAT 220 Acc GAT 512 CTC Val 1. Z ATC Met P PC Val A GCT 9,50 5 3 GAT Asp CTT Val 600 Ala 155 r g 250 re d GGC G1y 235 8 8 2 2 E ig **Arg** TCT Ser E & Fred Phe GPG GLn AGC ABD 83 8 2 걸 GTT Val Ct7 90 ct E Z 8 g 경착당 818 250 Asn ABD 185 GAG G1u CAC 66C 61y F 52 6 th 53 15 S A 65 15 g 265 265 265 Leu 2523 ŧŝ 950 5,50 g g A H AGC Ser 1 833 S S JCC Ser £ 3 Ala 120 25 gi Val Val Va I 21,48 23 23 ម្ពុ ខ្ល сув Авр Ala Ala 135 135 135 ATC Met 55 Ser 돢 7GC Cy3 E 3 GAS GLu AAA Lys EFE CTT Pr Pr 215 g g 23 싫 že g 480 528 555 22 801 240 336 96 144 192 288 384 432 IGT GAT TTA CGG GIC CIG ICT ANA CTG CTG CGC GAC ICT CAC GIG CTG Ser P. G Eya Eya ATG Wet 75 925 ដ្ឋ CAT ABP Val GCC Ala 155 A CCA Val AAG CTAGARARA CCAAGGAGGT AATAAATA ATG AAA AGT CCT GCA CCA CCT (
Het Lys Ser Pro Ala Pro Pro 15
-2 E 3 Pro age Thr GTG 5538 AGC AAC Lys CGG Arg Thr 8543 2 63 Fe 6 CAC H13 Arg Pa 23 cke clu GCA Gly ម្ពុទ្ធ CAG Gln 105 c cs 23 Çe: TGC GTC Cys Val CAG C1u 12 kg SAC Asp AGC 534 GAG G1u TCC Ser 53 Arg A Leu 535 32 525 E G GTC ATC Met 55 re d TCA Ser ALa Ala a H (xi) SEQUENCE DESCRIPTION: SEQ ID NO:14: 25.35 £ 3 15 E G 955 걸 E 3 85. ម្ពុជ្ញ ដូខ្ល ŧŝ Val Met Ęŝ 뫍 ES 35 75 85 85 5 E E G (vi) ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens Val S H 計员 Pro MOLECULE TYPE: synthetic DNA (i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 1043 base pai
(B) TYPE: nucleic acid
(C) STRANDENESS: double
(D) TOPOLOGY: linear Lys Lys 14 960 ខ្មីដូ TCC Ser 뀱 523 TGA CAT HIS ម្ល ដូដូ Val INFORMATION FOR SEQ ID NO:14: Ser Ser 5 t Gre 2 2 553 52 CAS 감 95.5 F 38 F g Ę 7 7 8 8 25.4 GL GLy 5 5 8 8 6 8 Program AGC Ser 130 99 P. S. 88 GAC ដូដូ 52. Val GTA Val 145 F. 63 CAC 61.9 G1.4 623 613 65 15 33 15 33 Val i di 53 Ser 160 160 re Se d S C C 53 TTC Phe 53 ŧş S S 23 358 36.73 CJ.c Thr. ATC 11e ATC Het HTT 11e GAG Gln 5 g 3 GAG Glu AGC 623 617 95 95 53 thr Glo Asp 1 PC SCC Nta 5 3 TCT 5 H 51 95 g Ę Š ម្លដ្ឋន Eag 34 re Te (11) F H AGT 225 GAC ABP A S Ęŝ 65.4 61.4 AAT Aen 125 i i 2 20 6 E 3 146 E S 55 Val មិនីខ 3 5 5 53 A GG 250

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1012 1043 916 964 CCG ACC CCG ACC CCG CTG CTG AAC ACC ACC TAI ACC CAI ACC CAG Pro Thr Pro Thr Ser Pro Leu Leu Asn Thr Ser Tyr Thr His Ser Gln 320 CCG GTG GTT CAG CTG CAG CTG CCG GAT CCG AGC GCG CCG ACC Pro Val Val Gla Leu Bis Pro Leu Leu Pro Asp Pro Ser Ala Pro Thr 395 CTG CCG ACC Leu Pro Thr CCG ACC Pro Thr 1 290 CCG ACC GGC CAG TAT ACC CTG TTT CCG CTG CCG Pro Thr Gly Gla fyr Thr Leu Phe Pro Leu Pro 280 AAC CTG AGC CAG GAA GGC TAATGAAGCT TGA ABn Leu Ser Gln Glu Gly 330

(2) INFORMATION FOR SEQ ID NO:15:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 45 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDRESS: single

(C) STRANDEDNESS: sin (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:15:

(ii) MOLECULE TYPE: Other nucleic acid

AACTGGAAGA ATTCGCGGC GCAGGAATTT TITITITIT TITIT

(2) INFORMATION FOR SEQ ID NO:16:

(1) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 11 base pairs
(B) TYPE: nucleic acid
(C) STRANDDNESS: single
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(x1) SEQUENCE DESCRIPTION: SEQ ID NO:16:

NATTCGGCAC GAG

13

(2) INFORMATION FOR SEQ ID NO:17:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 9 base pairs
(B) TYPE: nucleic acid
(C) STRANNEDNESS: annole
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:17:

CTCCTCCCG

(2) INFORMATION FOR SEQ ID NO:18:

(1) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 12 amino acids
(B) TYPE: amino acid
(C) STRANDEDNESS:
(D) TOPPOLOGY: linear

(11) MOLECULE TYPE: peptide

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ile Pro Val Pro Pro Ala Cys Asp Pro Arg Leu Leu (x1) SEQUENCE DESCRIPTION: SEQ ID NO:18:

(2) INFORMATION FOR SEQ ID NO:19:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 12 amino acids
(B) TRPS: amino acid
(C) STRANDENRSS:
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: peptide

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:19:

Thr Pro Val Pro Pro Ala Cys Asp Pro Arg Leu Leu

(2) INFORMATION FOR SEQ ID NO:20:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 12 amino acids
(B) TYPE: amino acid
(C) STRANDEDRES:
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: peptide

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:20:

Ser Pro Val Pro Pro Ala Cys Asp Pro Arg Leu Leu 1

(2) INPORMATION FOR SEQ ID NO:21:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 17 base palrs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: single
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(1x) FEATURE:

(B) NAME/KEY: modified base (B) LOCATION: 12 & 15 (D) OTHER INFORMATION: /mod base= i

(x1) SEQUENCE DESCRIPTION: SEQ ID NO:21:

ACCCTGGGG CNGANGA

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(2) INFORMATION FOR SEQ ID NO:22:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 17 base pairs
(B) TYPE: nucleic acid
(C) SYRANDEDRESS: aingle
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(ix) FEATURE:
 (A) NAME/KEY: modified base
 (B) LOCATION: 12 & 15

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er information	S DESCRIPTION: SEQ
(D) OTHER	ğ
9	SEQUENCI
	(x)

(2) INFORMATION FOR SEQ ID NO:23:

ACACTAGGAT CHAMMAA

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(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 17 base pair (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(ix) FEATURE:

(A) NAME/KEY: modified base (B) LOCATION: 12 & 15 (D) OTHER INFORMATION: /mod_base= i

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:23:

ACCCTGGGTG CNGANGA

11

(2) INFORMATION FOR SEQ ID NO:24:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 17 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(ix) FEATURE:

(A) NAME/KEY: modified base
(B) LOCATION: 12 & 15
(D) OTHER INFORMATION: /mod_base= i

(*1) SEQUENCE DESCRIPTION: SEQ ID NO:24:

ACCCTGGGCG CNGANGA

13

(2) INFORMATION FOR SEQ ID NO:25:

(1) SEQUENCE CHARACTERISTICS

(A) LENGTH: 17 base pair: (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

.(x!) SEQUENCE DESCRIPTION: SEQ ID NO:25:

GGGGGCGA CCTGGG

17

(2) INFORMATION FOR SEQ ID NO:26:

(A) LENGTH: 17 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (i) SEQUENCE CHARACTERISTICS:

(ii) MOLECULE TYPE: Other nucleic acid

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO:26:

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GCACCACGAA CACTAGG

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(2) INFORMATION FOR SEQ ID NO:27:

(A) LENGTH: 17 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (i) SEQUENCE CHARACTERISTICS:

(ii) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:27:

11

(2) INFORMATION FOR SEQ ID NO:28:

GETGETCETA CCCTGGG

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 17 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:28:

(2) INFORMATION FOR SEQ ID NO:29:

GCCGCCGCA CGCTGGG

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(x1) SEQUENCE DESCRIPTION: SEQ ID NO:29:

(2) INFORMATION FOR SEQ ID NO:30:

GGATCTIGGT TCATTCTCAA G

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 20 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: single
(D) TOPOLOGY: linear

(11) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:30:

CCTCAGACAG TGGTTCAAAG

(i) SEQUENCE CHARACTERISTICS:

(2) INFORMATION FOR SEQ ID NO:31:

(A) LENCTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear

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(ii) MOLECULE TYPE: Other nucleic acid

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5 71 7 (xi) SEQUENCE DESCRIPTION: SEQ ID NO.31: (xi) SEQUENCE DESCRIPTION: SEQ ID NO:34: (xi) SEQUENCE DESCRIPTION: SEQ ID NO:32: (xi) SEQUENCE DESCRIPTION: SEQ ID NO:33: (ii) MOLECULE TYPE: Other nucleic acid (ii) MOLECULE TYPE: Other nucleic acid (11) MOLECULE TYPE: Other nucleic acid (i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 21 base pairs
(B) TYPE: nucleic acid
(C) STRANDENESS: single
(D) TOPOLOGY: linear (i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 21 base pairs
(B) TYPE: nucleic acid
(C) STRANDENESS: shole
(D) TOPOLOGY: linear (i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 21 base pair:
(B) TYPE: nucleic acid
(C) STRANDEDNESS: single (C) STRANDEDNESS: si. (D) TOPOLOGY: linear (2) INPORMATION FOR SEQ ID NO:32: (2) INFORMATION FOR SEQ ID NO:33: (2) INFORMATION FOR SEQ ID NO:34: TGAAGTTCGT CTCCAACAAT C CAGAGTTAGT CTTGCGGTGA G cererecere recenerate CCAGGGTGTA CCTGGGTCCT

(ii) MOLECULE TYPE: Other nucleic acid (i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 21 base pairs
(B) TYPE: nucleic acid
(C) SYRANDENESS: aingle
(D) TOPOLOGY: linear

(2) INPORMATION FOR SEQ ID NO:35:

(*i) SEQUENCE DESCRIPTION: SEQ ID NO:35:

ATGGAGCTGA CTGATTTGCT C

(2) INFORMATION FOR SEQ ID NO:36:

7

(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid

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(C) STRANDEDNESS: single (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(x1) SEQUENCE DESCRIPTION: SEQ ID NO:36:

TGAAGTICGI CICCAACAAI C

7

(2) INPORMATION FOR SEQ ID NO:37:

(i) SEQUENCE CHARACTERISTICS:
(A) IENGTH: 21 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: angle
(D) TOPOLOGY: linear

(11) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:37:

TTGTGACCIC CGAGICCICA G

(2) INFORMATION FOR SEQ ID NO:38:

(1) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 21 base pai
(B) TYPE: nuclaic acid
(C) STRANDEDNESS: singl
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:38:

(2) INFORMATION FOR SEQ ID NO:39: TGACCCAGAG GGTGGACCCT C

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(*1) SEQUENCE DESCRIPTION: SEQ ID NO:39:

AGCAGAACCT CTCTAGTCCT C

(2) INFORMATION FOR SEQ ID NO:40:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single

(C) STRANDEDNESS: Sir (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:40:

ACACTGAACG AGCTCCCAAA C

(2) INFORMATION FOR SEQ ID NO:41:

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(A) LENGTH: 21 base pa (B) TYPE: nucleic acid (C) STRANDEDNESS: sing (D) TOPOLOGY: linear (i) SEQUENCE CHARACTERISTICS:

(ii) NOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:41:

(2) INFORMATION FOR SEQ ID NO:42: AACTACTGGC TCTGGGCTTC T

7

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 21 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: single
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(*i) SEQUENCE DESCRIPTION: SEQ ID NO:42:

AGGGATTCAG AGCCAAGATT C

7

(2) INFORMATION FOR SEQ ID NO:43:

(A) LENCTH: 31 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (1) SEQUENCE CHARACTERISTICS:

(ii) MOLECULE TYPE: Other nucleic acid

(*i) SEQUENCE DESCRIPTION: SEQ ID NO:43:

TAGCGGCCGC ITTITITIT ITTITIGGG G

(1) SEQUENCE CHARACTERISTICS: (2) INFORMATION FOR SEQ ID NO:44:

(A) LENGTH: 31 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(*i) SEQUENCE DESCRIPTION: SEQ ID NO:44:

TAGCGGCCGC ITTITITIT ITTITITANA A

(2) INFORMATION FOR SEQ ID NO:45:

(1) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 31 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(*i) SEQUENCE DESCRIPTION: SEQ ID NO:45:

TAGGGGGGG TITITITIT ITITITICE :

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(2) INPORMATION FOR SEQ ID NO:46:

(1) SEQUENCE CHARACTERISTICS: (A) LENGTH: 31 DASS E

(A) LENGTH: 31 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(x1) SEQUENCE DESCRIPTION: SEQ ID NO:46:

TAGGGGGGG TITITITI TITITIGGG C

33

(2) INFORMATION FOR SEQ ID NO:47:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 21 base pairs
(B) TYPE: nuclaic acid
(C) STRANDEDNESS: aingle
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:47:

TAGGGGGGG TITITITI I

(2) INFORMATION FOR SEQ ID NO:48:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 21 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: single
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(x1) SEQUENCE DESCRIPTION: SEQ ID NO:48:

TTGTGACCTC CGAGTCCTCA G

(2) INFORMATION FOR SEQ ID NO:49:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 21 base pai.
(B) TYPE: nucleic acid
(C) STRANDEDNESS: single
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:49:

AGGATGGGTT GGGGAAGGAG A

(2) INFORMATION FOR SEQ ID NO:50:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 21 base pairs
(B) TYPE: mulaic acid
(C) STRANDENESS: single
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:50:

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TGTGACCTC GGAGTCCTCA G	2) INFORMATION FOR SEQ ID NO:51:	(1) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDENESS: single
rereacere ce	2) INPORMATI	odas (1).

(11) MOLECULE TYPE: Other nucleic acid

(*i) SEQUENCE DESCRIPTION: SEQ ID NO:51:

AGGGAAGAGC GTATACTGTC C

21

(i) SEQUENCE CHARACTERISTICS:

(2) INFORMATION FOR SEQ ID NO:52:

(A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:52:

GCCAGCCAG ACACCCGGC C

(2) INFORMATION FOR SEQ ID NO:53:

(1) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 21 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: single
(D) TOPOLOGY: linest

(ii) MOLECULE TYPE: Other nucleic acid

(*i) SEQUENCE DESCRIPTION: SEQ ID NO:53:

ATGGGAGTCA CGAAGCAGTT T

7

(2) INFORMATION FOR SEQ ID NO:54:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(*i) SEQUÈNCE DESCRIPTION: SEQ ID NO:54:

(2) INFORMATION FOR SEQ ID NO:55:

TGCGTTTCCT GAIGCTTGIA G

7

(1) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 21 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: aingle
(D) TOPOLOGY: linear

(11) MOLECULE TYPE: Other nucleic acid

(x1) SEQUENCE DESCRIPTION: SEQ ID NO:55: AACCTTACCC TTCCTGAGAC A

7

(2) INFORMATION FOR SEQ ID NO:56:

(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 30 base

(D) TOPOLOGY: linear

(11) MOLECULE TYPE; Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:56:

TITGAAITCG GCCAGCCAGA CACCCCGGCC

8

(2) INFORMATION FOR SEQ ID NO:57:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 39 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: annyle
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:57:

THEGGGCCG CICAINAGCY GGGACAGCY GTGGTGGGT

(2) INFORMATION FOR SEQ ID NO:58:

(i) SEQUENCE CHANACTERISTICS:
(A) LENGTH: 42 base pairs
(B) TYPE: nucleic acid
(C) STRANDENNESS: single
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:58:

TITGCGGCCG CICAITACAG IGIGAGGACT AGAGAGGITC TG

42

(2) INFORMATION FOR SEQ ID NO:59:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 42 base pairs
(B) TYPE: nucleic acid
(C) STRANDENESS: single
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(*1) SEQUENCE DESCRIPTION: SEQ ID NO:59:

TTTGGGGGG CTCATTAICT GGCTGAGGCA GTGAAGTTTG TC

42

(2) INFORMATION FOR SEQ ID NO:60:

(1) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 42 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO:60: TITGGGGCG CICATIACAG ACCAGGAAIC INGGCICTGAA (11) MOLECULE TYPE: Other nucleic acid (2) INFORMATION FOR SEQ ID NO:61:

42

(1) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 10 base pairs
(B) TYPE: nucleic acid
(C) STRANDENESS: single
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(*i) SEQUENCE DESCRIPTION: SEQ ID NO:61:

(2) INFORMATION FOR SEQ ID NO: 62:

TITGAATICG GCCAGCCAGA CACCCGGGC

8

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 41 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(*i) SEQUENCE DESCRIPTION: SEQ ID NO:62:

TITGGGGCG CICATIAGAG GGTGGACCCI CCIACAAGCA I

4

(2) INFORMATION FOR SEQ ID NO:63:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 40 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(*i) SEQUENCE DESCRIPTION: SEQ ID NO:63:

TTTGGGGCG CTCATTAGCA GAGGOTGGAC CCTCCTACAA (2) INPORMATION FOR SEQ ID NO:64:

6

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 38 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear

(11) MOLECULE TYPE: Other nucleic acid

(*i) SEQUENCE DESCRIPTION: SEQ ID NO:64: TITGCGGCCG CICAITACCI GACGCAGAGG GIGGACCC

(2) INFORMATION FOR SEQ ID NO:65:

38

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 18 base pairs
(B) TYPE: nucleic acid

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(C) STRANDEDNESS: single (D) TOPOLOGY: linear

(11) MOLECULE TYPE: Other nucleic acid

(*1) SEQUENCE DESCRIPTION: SEQ ID NO:65:

TITGCGCCC CICAIIACCC CCIGACGCAG AGGGIGGA

38

(2) INFORMATION FOR SEQ ID NO:66:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 18 base pairs
(B) TYPE: nucleic acid
(C) STRANDENESS: single
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:66:

TITGCGGCCG CICATIAGGC CCGCCIGACG CAGAGGGI

38

(2) INPORMATION FOR SEQ ID NO:67:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 38 base pairs
(B) TYPE: nucleic acid
(C) STRANDENESS: single
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:67: TITGCGCCC CICALIAIGG GCCCCCCTG ACGCAGAG

38

(2) INFORMATION FOR SEQ ID NO:68:

(A) LENGTH: 38 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (i) SEQUENCE CHARACTERISTICS:

(ii) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:68:

TITGCGGCCG CICAITAGGG IGGGGCCCGC CIGACGCA

38

(2) INFORMATION FOR SEQ ID NO:69:

(A) LENGTH: 30 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (i) SEQUENCE CHARACTERISTICS:

(11) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:69:

TITGAAITCG GCCAGCCAGA CACCCCGGCC

(2) INPORMATION FOR SEQ ID NO:70:

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TYPE: nucleic acid STRANDEDNESS: single (i) SEQUENCE CHARACTERISTICS: £200

TOPOLOGY: linear

(*1) SEQUENCE DESCRIPTION: SEQ ID NO:70:

(11) MOLECULE TYPE: Other nucleic acid

TITECGCCC CTCATTAITC GIGIAICCTC IICAGGIAIC

41

(2) INFORMATION FOR SEQ ID NO:71:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 39 base pairs
(B) TYPE: nucleic acid
(C) STRANDENESS: single
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(*1) SEQUENCE DESCRIPTION: SEQ ID NO:71:

ITTCCGCCC CICATTAGCT GGGACACCT GIGGIGGGI

33

(2) INFORMATION FOR SEQ ID NO:72:

(i) SEQUENCE CHARACTERISTICS:

LENGTH: 50 base pairs TYPE: nucleic acid STRANDEDNESS: single (A) LENGTH: 50 base pa (B) TYPE: nucleic acid (C) STRANDEDNESS: sing (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:72:

AGTAAACTGC IICGIGACTC CCAIGICCII CACAGCAGAC IGAGCCAGIG

(2) INFORMATION FOR SEQ ID NO:73:

(1) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 49 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:73:

CATGGGAGTC ACGAAGCAGT TTACTGGACA GCGTTAGCCT TGCAGTTAG

(2) INFORMATION FOR SEQ ID NO:74:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 49 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear

(*i) SEQUENCE DESCRIPTION: SEQ ID NO:74:

(ii) MOLECULE TYPE: Other nucleic acid

TGTGACCTCC GAGTCCTCAG TAAACTGCTT CGTGACTCCC ATGTCCTTC

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(2) INFORMATION FOR SEQ ID NO:75:

(1) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 48 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear

(11) MOLECULE TYPE: Other nucleic acid

(*i) SEQUENCE DESCRIPTION: SEQ ID NO:75:

TTTACTGAGG ACTCGGAGGT CACAGGACAG CGTTAGCCTT GCAGTTAG

48

(2) INFORMATION FOR SEQ ID NO:76:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 49 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(x1) SEQUENCE DESCRIPTION: SEQ ID NO:76:

GACCTCCGAG ICCTCAGIAA ACTGCTICGI GACTCCCAIG ICCTICACA

(2) INFORMATION FOR SEQ ID NO:77:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 48 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:77:

CAGITIACIG AGGACICGGA GGICGGACAG CGIIAGCCII GCAGIIAG

(2) INFORMATION FOR SEQ ID NO:78:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 21 base pairs (B) TYPE: nuclaic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:78:

(2) INFORMATION FOR SEQ ID NO:79:

TTGTGACCTC CGAGTCCTCA G

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 21 base pairs
(B) TYPE: nucleic acid
(C) STRANDENESS: single
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:79:

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CAGGIATCCG GGGATTIGGI C

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(2) INFORMATION FOR SEQ ID NO:80:

(1) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 21 base palre
(B) TYPE: nucleic acid
(C) STRANDEDNESS: single
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:80:

TECCTITCCI GAIGCITGIA G

57

(2) INFORMATION FOR SEQ ID NO:81:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 35 base pairs
(B) TYPE: nucleac acid
(C) STRANDENESS: single
(D) TOPOLOGY: lines

(ii) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:81:

GAGAGAGGGG COGCTTACCC TTCCTGAGAC AGATT

32

(2) INFORMATION FOR SEQ ID NO:82:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 6 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOCY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(*i) SEQUENCE DESCRIPTION: SEQ ID NO:82:

GAGAGA

(2) INFORMATION FOR SEQ ID NO:83:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 70 base pairs
(B) TYPE: nucleic acid
(C) STRANDENESS: single
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(*1) SEQUENCE DESCRIPTION: SEQ ID NO:83:

8 ctagaaaaa ccaaggaggt aataaataat gagcooggct cogccagctt gtgaccttcg TGITCICICI

(2) INPORMATION FOR SEQ ID NO:84:

(i) SEQUENCE CHANACTERISTICS:
(A) IZEGTH: 72 base pairs
(B) TYPE: nucleic acid
(C) STRANDENESS: single
(D) TOPOLOGY: lines

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TYPE:	
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(*i) SEQUENCE DESCRIPTION: SEQ ID NO:84:

ŝ CAGTITAGAC AGAACACGAA GGICACAAGC TGGCGGAGCC GGGCTCAITA ITTAITACCT CCTTGGTTTT IT

(2) INFORMATION FOR SEQ ID NO:85:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 69 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:85:

AAACIGCIIC GOGACICICA CGIGCIGCAC ICICGICIGI CCCAGIGCCC GGAAGIICAC 60

(2) INFORMATION FOR SEQ ID NO:86:

CCGCTGCCG

(i) SEQUENCE CHAIACTERISTICS:
(A) LENGTH: 69 base pairs
(B) TYPE: nucleic acid
(C) STRANDENESS: single
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(xì) SEQUENCE DESCRIPTION: SEQ ID NO:86:

CGGGGICGGC AGCGGGTGAA CTTCCGGGCA CTGGGACAGA CGAGAGIGCA GCACGTGAGA 60 GTCCCGAAG

(2) INFORMATION FOR SEQ ID NO:87:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 69 base pairs
(B) TYPE: nucleic acid
(C) STRANDENESS: single
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(*1) SEQUENCE DESCRIPTION: SEQ ID NO:87:

ACCCCGGITC TGCITCCGCC TGTCGACTIC TCCCTGGGTG AATGGAAAAC CCAGATGGAA 60 GAGACCAAA

(2) INFORMATION FOR SEQ ID NO:88:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 69 base pairs
(B) TYPE: nucleic acid
(C) STRANDENESS: single
(D) TOPOLOGY: lines

(11) MOLECULE TYPE: Other nucleic acid

(*1) SEQUENCE DESCRIPTION: SEQ ID NO:88:

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TGAGCITIG GICTCTICCA TCTGGGTTTI CCATTCACCC AGGGAGAAGI CGACAGCGG 60		
AGGGAGAAGT		
CCATTCACCC		
TCTGGGTTTT		
GTCTCTTCCA		
TGAGCTTTC	AGCAGAAC	

(2) INPORMATION FOR SEQ ID NO:89:

(1) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 69 base pairs (B) TYPE: nucleic acid (C) STANDEDNESS: single (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:89:

GCTCAGGACA ICCIGGGIGG AGIAACICIG CTICIGGAAG GCGIIAIGGC IGCACGIGGC 60 CAGCTTGGC

(2) INPORMATION FOR SEQ ID NO:90:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 69 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: single
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(*i) SEQUENCE DESCRIPTION: SEQ ID NO:90;

8 GGICGGGCCA AGCTGGCCAC GIGCAGCCAI AACGCCTICC AGAAGCAGAG ITACTGCACC CAGGATGIC

(2) INFORMATION FOR SEQ ID NO:91:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 69 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(*1) SEQUENCE DESCRIPTION: SEQ ID NO:91:

CCGACCTGCC TOTCTTCCCT GCTTGGCCAG CTGTCTGGCC AGGTTCGTCT GCTGCTCGGC 60 GCTCTGCAG

(2) INFORMATION FOR SEQ ID NO:92:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 66 base pairs
(B) TYPE: nucleic seid
(C) STRANDENESS: single
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:92:

Chgrgretge agagegeega gergregare etggeergae agetggeera gerggarga 60 CAGGCA

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(1) SEQUENCE CHARACTERISTICS:

LENGTH: 70 base pairs TYPE: nucleic acid STRANDEDNESS: single E ê

(C) STRANDEDNESS: SL (D) TOPOLOGY: linear

(11) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:93:

8 TCTCTGCTTG GCACCCAGGT GCCGCCACAG GGCCGTACCA CTGCTCACAA GGATCCGAAC GCTATCTTCC

(2) INFORMATION FOR SEQ ID NO:94:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 70 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single

(C) STRANDEDNESS: 81 (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:94:

ဇ္ AAGACAGGAA GAIAGCGIIC GGAICCIIGI GAGCAGIGGI ACGGCCCIGI GGCGGCAGCI GGGTGCCAAG

6

(2) INFORMATION FOR SEQ ID NO:95:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(*i) SEQUENCE DESCRIPTION: SEQ ID NO:95:

GGAGGAGACC AAGGCACAGG A

(2) INFORMATION FOR SEQ ID NO:96:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 30 base pairs
(B) TYPE: nucleic acid
(C) STRANDENESS: single
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(*i) SEQUENCE DESCRIPTION: SEQ ID NO:96:

(2) INFORMATION FOR SEQ ID NO:97:

COGGNATICT TACCCTICCT GAGACAGAIT

8

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 8 base pair (B) TYPE: nucleic acid (C) STRANDEDNESS: sing] (D) TOPOLOGY: linear

TYPE: nucleic acid STRANDEDNESS: single TOPOLOGY: linear

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TYPE:
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(x1) SEQUENCE DESCRIPTION: SEQ ID NO:97:

CINATIGAG

(2) INFORMATION POR SEQ ID NO:98:

- (i) SEQUENCE
- (A) LENGTH: 16 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Other nucleic acid
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:98:

AATTCTCATT AGAGCT

(2) INFORMATION FOR SEQ ID NO:99:

- (1) SEQUENCE CHARACTERISTICS:
- (A) LENGTH: 8 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear

- (II) MOLECULE TYPE: Other nucleic acid
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:99:

CTAATGAG

(2) INFORMATION FOR SEQ ID NO: 100:

- (1) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 16 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear
- (11) MOLECULE TYPE: Other nucleic acid
- (*i) SEQUENCE DESCRIPTION: SEQ ID NO:100;

AATTCTCATT AGAGCT

(2) INFORMATION FOR SEQ ID NO:101:

16

- (A) LENGTH: 25 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (i) SEQUENCE CHARACTERISTICS:
- (11) MOLECULE TYPE: Other nucleic acid
- (*t) SEQUENCE DESCRIPTION: SEQ ID NO:101:

ATCGCCGGCT CCGCCAGCTT GTGAC

25

(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 30 base paire (B) TYPE: nucleic acid

(2) INFORMATION FOR SEQ ID NO:102:

- (C) STRANDEDNESS: single (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Other nucleic acid
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:102:

GCCGAATICT CATTAGAGCT CGTTCAGIGT

8

- (2) INFORMATION FOR SEQ ID NO: 103:
- (A) LENGTH: 42 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (i) SEQUENCE CHARACTERISTICS:
- (ii) MOLECULE TYPE: Other nucleic acid
- (x1) SEQUENCE DESCRIPTION: SEQ ID NO:103:
- GAICCGAACG CIAICITCCT GICTITCCAG CACCIGCICC GI

42

- (2) INFORMATION FOR SEQ ID NO:104:
- (i) SEQUENCE CHARACTERISTICS:
 (A) LENGTH: 44 base pairs
 (B) TYPE: nucleic acid
 (C) STRANDENESS: single
 (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Other nucleic acid
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:104:

TITGCCACGC AGCAGGTGCT GGAAAGACAG GAAGAIAGCG ITCG

- (2) INFORMATION FOR SEQ ID NO:105:
- (A) LENGTH: 42 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (i) SEQUENCE CHARACTERISTICS:
- (ii) MOLECULE TYPE: Other nucleic acid
- GGCAAAGIIC GITICCIGAI GCIGGIIGGC GGIICIACGC IG (xi) SEQUENCE DESCRIPTION: SEQ ID NO:105:

42

(2) INFORMATION FOR SEQ ID NO: 106:

- (i) SEQUENCE CHARACTERISTICS:
 (A) LENGTH: 41 base pairs
 (B) TYPE: nucleic acid
 (C) STRANDENESS: angle
 (D) TOPOLOGY: linesr

- (ii) MOLECULE TYPE: Other nucleic acid
- (*1) SEQUENCE DESCRIPTION: SEQ ID NO:106: ACGCACAGGG TAGAACCGCC AACCAGCATC AGGAAACGAA C

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(2) INFORMATION FOR SEQ ID NO: 107:

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(A) LENGTH: 46 base pairs (B) TYPE: nucleic acid (C) STRANDENPES: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: Other nucleic acid	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:107:	
TGCGITCGIC GGGGGCGGC AACCACTGCI GITCCGICIT AAIGAA	46
(2) INFORMATION FOR SEQ ID NO:108:	:
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 45 base pairs (B) TYPE: nucleic acid (C) STRANDENESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: Other nucleic acid	-
(*1) SEQUENCE DESCRIPTION: SEQ ID NO:108:	
AGCITICATI AAGACGGAAC AGCAGIGGII GGCGGCGCC GACGA	46
(2) INFORMATION FOR SEQ ID NO:109:	
(i) SEQUENCE CHANACTERISTICS; (A) LENGTH: 24 base pairs (B) TYPE: nuclealc acid (C) STRANDENESS: single (D) TOPOLOGY: lines	
(ii) MOLECULE TYPE: Other nucleic acid	
(*1) SEQUENCE DESCRIPTION: SEQ ID NO:109:	
AAGGATCCGA ACGCTATCTT CCTC	
(2) INFORMATION FOR SEQ ID NO:110;	5
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 24 base paics (B) TYPE: nucleic acid (C) STRANDEDRESS: single (D) TOPOLOCY: lines:	
(ii) MOLECULE TYPE: Other nucleic acid	
(*i) SEQUENCE DESCRIPTION: SEQ ID NO:110:	
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(2) INPORMATION FOR SEQ ID NO:111;	57
(1) SEQUENCE CHARACTERISTICS: (A) LENGTH: 46 base pairs (B) TYPE: nucleace acid (C) STRANDEDERSS: single (D) TOPOLOCY: linear	
(ii) MOLECULE TYPE: Other nucleic acid	
(*1) SEQUENCE DESCRIPTION: SEQ ID NO:111:	
CTAGAAAAA CCAAGGAGGI AATAAATAAI GAAAGGACCT GTACCA	. 46

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- (1) SEQUENCE CHARACTERISTICS:
- (A) LENGTH: 46 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Other nucleic acid
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:112:
- CAGGIGGIAC AGGIGCITIC AITAITIAIT ACCICCIIGG ITITIT

46

- (2) INFORMATION FOR SEQ ID NO:113:
- (i) SEQUENCE CHARACTERISTICS:
 (A) LENGTH: 3B base pairs
 (B) TYPE: nucleic acid
 (C) STRANDENESS: single
 (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Other nucleic acid
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:113:
- CCTGCATGTG ATTTACGGGT CCTGTCTAAA CTGCTGCG

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- (2) INFORMATION FOR SEQ ID NO:114:
- (i) SEQUENCE CHARACTERISTICS:
 (A) LENGTH: 34 base pairs
 (B) TYPE: nucleic acid
 (C) STRANDENESS: single
 (D) TOPOLOGY: linear

- (11) MOLECULE TYPE: Other nucleic acid
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:114:
- GGCAGCAGIT TAGACAGGAC COGTAAAICA CAIG

34

- (2) INFORMATION FOR SEQ ID NO:115:
- (i) SEQUENCE CHARACTERISTICS:
 (A) IERGTH: 84 base pairs
 (B) TYPE: nucleic acid
 (C) STRANDENESS: double
 (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Other nucleic acid
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:115:

CTAGAAAAA CCAAGGAGGI AAINAAIAAI GAAAGCACCI GIACCACCIG CAIGIGAIII 60 ACGGGTCCTG TCTAAACTGC TGCG

- (2) INFORMATION FOR SEQ ID NO:116:
- (i) SEQUENCE CHARACTERISTICS:
 (A) LENGTH: 20 amino acids (B) TYPS: amino acid (C) STRANDEDNESS:
 (D) TOPOLOGY: linear

- (ii) MOLECULE TYPE: peptide

(*i) SEQUENCE DESCRIPTION: SEQ ID NO:116:

Pro Arg Leu Leu Asn Lys Leu Leu Arg Asp Ser Tyr Leu Leu His Arg $_{\rm 1}$

Arg Leu Ser Gln 20

(2) INFORMATION FOR SEQ ID NO:117:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 21 amino acids
(B) TYPE: amino acid
(C) STRANDEDNESS:
(D) TOPOLOGY: linear

(11) MOLECULE TYPE: peptide

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:117:

Phe Ser Leu Gly Glu Trp Lys Thr Gln Thr Glu Gln Ser Lys Ala Gln $_{\rm 1}$

Asp Ile Leu Gly Ala 20

(2) INFORMATION FOR SEQ ID NO:118:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 18 amino acids
(B) TTPE: amino acid
(C) STRANDEDNESS:
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: peptide

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:118:

Ser Arg Thr Ser Gin Leu Leu Thr Leu Asn Lys Phe Pro Asn Arg Leu $_{\rm 1}$ $_{\rm 1}$

Leu Asp

(2) INFORMATION FOR SEQ ID NO:119:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 21 amino acids
(B) TYPE: amino acid
(C) STRANDEDNESS:
(D) TOPOLOGY: linear

(11) MOLECULE TYPE: peptide

(*1) SEQUENCE DESCRIPTION: SEQ ID NO:119:

Asp Leu Arg Val Leu Ser Lys Leu Leu Arg Asp Ser His Val Leu His $_{\rm 1}$

Ser Arg Leu Ser Gln 20

(2) INFORMATION FOR SEQ ID NO:120:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 16 amino acids
(B) TYPE: amino acid
(C) STRANDEDNESS:

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(D) TOPOLOGY: linear

(11) HOLECULE TYPE: peptide

(x1) SEQUENCE DESCRIPTION: SEQ ID NO:120:

Ser Leu Gly Glu Trp Lys Thr Gln Met Glu Glu Thr Lys Ala Gln Asp

(2) INFORMATION FOR SEQ ID NO:121:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 19 amino acid
(B) TTPE: amino acid
(C) STRANDEDNESS:
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: paptide

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:121:

Leu Gly Thr Gln Leu Pro Pro Gln Gly Arg Thr Thr Ala His Lys Asp 1 5 15

Pro Asn Ala

(2) INFORMATION FOR SEQ ID NO:122:

(1) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 19 amino acids (B) TYPS: amino acid (C) STRANDEDNESS:
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: peptide

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:122:

Asn Glu Leu Pro Asn Arg Thr Ser Gly Leu Leu Glu Thr Asn Phe Thr $_{\rm 1}$

Ala Ser Ala

(2) INFORMATION FOR SEQ ID NO:123:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 23 amino acids
(B) TYPE: amino acid
(C) STRANDEDNESS:
(D) TOPOLOGY: Linear

(11) MOLECULE TYPE: peptide

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:123:

Ser Leu Pro Pro Aan Leu Gln Pro Gly Tyr Ser Pro Ser Pro Thr His 1

Pro Pro Thr Gly Gln Tyr Thr 20

(2) INFORMATION FOR SEQ ID NO:124:

(A) LENGTH: 27 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: (1) SEQUENCE CHARACTERISTICS:

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(D) TOPOLOGY: linear

(11) MOLECULE TYPE: peptide

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:124:

Pro Ser Ala Pro Thr Pro Thr Ser Pro Leu Leu Asn Thr Ser 1

Tyr Thr His Ser Gin Asn Leu Ser Gin Glu Gly 20

(2) INFORMATION FOR SEQ ID NO:125:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 46 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(*!) SEQUENCE DESCRIPTION: SEQ ID NO:125:

CTAGARARA CCAAGGAGGT RATARATRAT GARATCTCCT GCACCA

46

(2) INFORMATION FOR SEQ ID NO:126:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 46 base pairs
(B) TYPE: nucleic acid
(C) STRANDEBNESS: single

(C) STRANDEDNESS: Bir (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:126: (ii) MOLECULE TYPE: Other nucleic acid

CAGGIGGIGC AGGAGAITIC AITAITIAIT ACCICCING IIIIII

(2) INFORMATION FOR SEQ ID NO:127:

(i) SEQUENCE CHAIACTERISTICS:
(A) LENGTH: 38 base pairs
(B) TYPE: nucleic acid
(C) STRANDENESS: single
(D) TOPOLOGY: linear

(11) MOLECULE TYPE: Other nucleic acid

(*1) SEQUENCE DESCRIPTION: SEQ ID NO:127:

CCTGCATGTG ATTTAGGGT CCTGTCTAAA CTGCTGCG (2) INFORMATION FOR SEQ ID NO:128:

38

(1) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 34 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(*1) SEQUENCE DESCRIPTION: SEQ ID NO:128:

CGCAGCAGIT TAGACAGGAC CCGIAAAICA CAIG

3

(2) INFORMATION FOR SEQ ID NO:129:

(i) SEQUENCE CHARACTERISTICS:

(B) TYPE: nucleic acid (C) STRANDEDNESS: double (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:129:

8 CTRGRADADA CCADGGRGGT DATADATANT GADATCTCCT GCACCACCTG CATGTGATTT

ACGGGTCCTG TCTAAACTGC TGCG

84

(2) INFORMATION FOR SEQ ID NO:130:

(i) SEQUENCE CHARACTERISTICS:
(A) LENCTH: 47 base pairs
(B) TYPE: nucleic acid
(C) STRANDENESS: single
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:130:

CTCCCGAACC GTACCAGCG CCTGCTCGAA ACCAACTTTA CCGCGAG

47

(2) INFORMATION FOR SEQ ID NO:131:

(1) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 47 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:131:

GGTAAAGTIG GITTCCAGCA GGCCGCTGGT ACGGTTCGGG AGCTCGT

47

(2) INFORMATION FOR SEQ ID NO:132:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 49 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: aingle
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:132:

COCCCGINC ACCCCNCC CCTGCTGAN AIGGCAGCAG GCTTICGI

(2) INFORMATION FOR SEQ ID NO:133:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 49 base pairs
(B) TYPE: nucleic acid
(C) STRANDENESS: single
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

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49 AGCCCTGCTG CCAITICAGC AGGCCGCTGC CGGTGGTACG CGCCTCGC (xi) SEQUENCE DESCRIPTION: SEQ ID NO:133: 319

(2) INPORMATION POR SEQ ID NO:134:

(1) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 50 base pairs
(B) TYPE: nucleac acid
(C) STRANDENESS: single
(D) TOPOLOGY: linear

(11) MOLECULE TYPE: Other nucleic acid

GOGARANTEC CEGGCCTGCT GAACCAGACC AGCCGTAGCC TGGATCAGAT (xi) SEQUENCE DESCRIPTION: SEQ ID NO:134:

20

(2) INFORMATION FOR SEQ ID NO:135:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 50 base pairs

TYPE: nucleic acid STRANDEDNESS: single

(C) STRANDEDNESS: Bir (D) TOPOLOGY: linear

(11) MOLECULE TYPE: Other nucleic acid

(x1) SEQUENCE DESCRIPTION: SEQ ID NO:135:

8 ATCCAGGCIA COGCIGGICI GGIICAGCAG GCCCGGGAII IICGCACGAA

(2) INFORMATION POR SEQ ID NO:136:

(1) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 47 base pairs
(B) TYPE: nucleic acid
(C) STRANDENESS: single
(D) TOPOLOGY: lines in

(11) MOLECULE TYPE: Other nucleic acid

(x1) SEQUENCE DESCRIPTION: SEQ. ID NO:136:

4 CCCGGGCTAT CTGAACCGTA TCCATGAACT GCTGAACGC ACCCGTG

(2) INFORMATION FOR SEQ ID NO:137:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 47 base pairs
(B) TYPE: nucleic acid
(C) STRANNEDNESS: aingle
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:137:

CTCCCCTTCA CCAGTTCATG CATACGGTTC AGATAGGGGG GGATCTG

(2) INFORMATION FOR SEQ ID NO:138:

(1) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 49 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear

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(ii) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:138:

49 econstruct sescocases cerescases rescoseses seatanteas

(2) INFORMATION FOR SEQ ID NO:139:

(1) SEQUENCE

(A) LENGTH: 49 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:139:

ATCCCCCCC CCCAGGGTGC CACGGTCGG GCCCGGAAAC AGGCCACGG

49

(2) INFORMATION FOR SEQ ID NO:140:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: SO base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:140:

20 ATCAGCTCTG GCACCAGCGA TACCGGCAGC CTGCCGCCGA ACCTGCAGCC

(2) INFORMATION FOR SEQ ID NO:141:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 50 base paire
(B) TYPE: nucleic acid
(C) STRANDEDNESS: aingle
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:141:

20 CAGGITCGGC GGCAGGTTGC GGTGCCAGAG CIGATATCCG

(2) INFORMATION FOR SEQ ID NO:142:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 51 base pairs
(B) TYPE: nucleic acid
(C) STRANDENNESS: single
(D) TOPOLOGY: linear

(11) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:142:

GGGCTATAGC CCGAGCCCGA CCCATCCGCC GACCGGCCAG TATACCCTGT T (2) INFORMATION FOR SEQ ID NO:143:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGIR: 51 base pairs
(B) TYPE: nucleic acid

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- (C) STRANDEDNESS: single (D) TOPOLOGY: linear
- (11) MOLECULE TYPE: Other nucleic acid
- (x1) SEQUENCE DESCRIPTION: SEQ ID NO:143:

ន GGTATACTICG COGGTCGGCG GATGGGCTGG GCTCGGGCTA TAGCCCGGCT G

- (2) INPORMATION FOR SEQ ID NO:144:
- (i) SEQUENCE CHARACTERISTICS:
 (A) LENGTH: 50 base pairs
 (B) TYRYE: nucleic acid
 (C) STRANDENESS: single
 (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Other nucleic acid
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:144:

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- (2) INFORMATION FOR SEQ ID NO:145:

- (1) SEQUENCE CHARACTERISTICS:
 (A) LENGTH: SO base pairs
 (B) TYPE: nuclaic acid
 (C) STRANDENESS: single
 (D) TOPOLOGY: linear
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:145: (ii) MOLECULE TYPE: Other nucleic acid
- င္တ GENTGCAGCT GAACCACCGC GGTCGCCAGG GTCGCCGCCA GCGGAAACAG
- (2) INFORMATION FOR SEQ ID NO:146:
- (1) SEQUENCE CHARACTERISTICS:
- (A) LENGTH: 51 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Other nucleic acid
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:146:

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- (2) INFORMATION FOR SEQ ID NO:147:
- (1) SEQUENCE CHARACTERISTICS:
 (A) LENGTH: 51 base pairs
 (B) TYPE: nucleic acid
 (C) STRANDENESS: single
 (D) TOPOLOGY: iinear

- (ii) MOLECULE TYPE: Other nucleic acid
- (x1) SEQUENCE DESCRIPTION: SEQ ID NO:147:

2 AGCAGCGGG TGGTCGGGGT CGGGTCGGC GCGCTCGGAT CCGGCAGCAG C

(2) INFORMATION FOR SEQ ID NO: 148:

322

- (A) LENGTH: 51 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (1) SEQUENCE CHARACTERISTICS:
- (1i) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:148:

CCAGCTATAC CCATAGCCAG AACCTGAGCC AGGAAGGCTA ATGAAGCTTG A

2

- (2) INFORMATION FOR SEQ ID NO:149:
- (i) SEQUENCE CHARACTERISTICS:
 (A) LENGTH: 51 base pairs
 (B) TYPE: nucleic acid
 (C) STRANDENESS: single
 (D) TOPOLOGY: linear

- (ii) MOLECULE TYPE: Other nucleic acid
- (*1) SEQUENCE DESCRIPTION: SEQ ID NO:149:

CTICATIAGC CTICCIGGCI CAGGITCIGG CIAIGGGIAI AGCIGGIGII C

- (2) INFORMATION FOR SEQ ID NO:150:
- (i) SEQUENCE CHARACTERISTICS:
- (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear
- (11) MOLECULE TYPE: Other nucleic acid
- (*1) SEQUENCE DESCRIPTION: SEQ ID NO:150:

ACGAGCTCCC GAACCGTACC A

7

- (2) INFORMATION FOR SEQ ID NO:151:
- (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (1) SEQUENCE CHARACTERISTICS:
- (ii) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:151:

CTGATATCCG GCGCCCCAG G

(2) INFORMATION FOR SEQ ID NO:152:

7

- (1) SEQUENCE CHARACTERISTICS:
 (A) LENGTH: 21 base pairs
 (B) TYPE: nucleic acid
 (C) STRANDENESS: single
 (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Other nucleic acid
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:152:

CGGATATCAG CTCTGGCACC A

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48 96 144 192 240 288 336 384 432 480 490 CCC GAT ATC Pro Asp Ile 80 CTG CAG CCC CTG CAT CCG ACC CAI AGC CAG AAC CIG AGC CAG GAA GGC TAA Thr His Ser Gin Asn Leu Ser Gin Giu Gly 145 CTG CTG AAC CAG ACC AGC CGT AGC CTG GAT CAG Leu Leu Asn Gln Thr Ser Arg Ser Leu Asp Gln 45 ខ្លួ AGC £5.57 83 A 251 77 ACC GCG 1 Thr Ala TAT ACC (Tyr Inc 1 CCG ACC AGC CCG (Pro Thr Ser Pro I GGG CGT ACC ACC GGC AGC GGC CTG AAA TGG CAG GGC TTT ALA Arg Thr Thr Gly Ser Gly Leu Leu Lys Trp Gln Gln Gly Phe. 25 AAC GGC ACC ABn Gly Thr : ACC CTG GGC GCG CT 7 Thr Leu Gly Ala P CTG CCG CCG AAC C Leu Pro Pro Asn L 90 CCG AAC CGI ACC AGC GGC CTG GTG GAA ACC AAC TIT Pro Asn Arg Thr Ser Gly Leu Leu Glu Thr Asn Phe 25.2 25 Gla AAC CGT ATC CAT GAA CTG CTG ? Asn Arg Ile His Glu Leu Leu ? 55 60 950 t s (xi) SEQUENCE DESCRIPTION: SEQ ID NO:154: (*1) SEQUENCE DESCRIPTION: SEQ ID NO:153: CCC ACC O CCG GTG G CCG ACC C (ii) MOLECULE TYPE: Other nucleic acid (ii) MOLECULE TYPE: Other nucleic acid (A) LENGTH: 490 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: double (D) TOPOLOGY: linear CCG AGC CGT CGC A CCG ACC C Pro Thr B ACC AGC GAT ACC GGC AGC Thr Ser Asp Thr Gly Ser 85 a i 19 P C C (2) INPORMATION FOR SEQ ID NO:153: (1) SEQUENCE CHARACTERISTICS: (2) INFORMATION FOR SEQ ID NO:154: TOPOLOGY: 11near (1) SEQUENCE CHARACTERISTICS: GCG CCG A Ala Pro 1 135 CAT His CCG ACC Pro Thr ACC CTG Thr Leu AGC TCAAGCTTCA TTAGCCTTCC T Pro Gly B AGC TAT A Ser Tyr 1 Pro Pro 1 GCG AAA ATC CCG GGC Ala Lys Ile Pro Gly 35 i G CCG AGC Pro Ser 100 S S €€0€ GAT (IN I GGC CTG TTT C Gly Leu Phe P 65 ATC CCG GGC 7 Ile Pro Gly 7 50 Ser Gly 1 GGC TAT AGC Gly Tyr Ser 1 CTG AAC ACC Leu Aan Thr 115 8 2 TGAAGCTTGA TTT CCG 250 23 Ser 23

(2) INFORMATION FOR SEQ ID NO:155:

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(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 24 base pairs
(B) TYPE: nucleic acid
(C) STRANDENESS: single
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(x1) SEQUENCE DESCRIPTION: SEQ ID NO:155:

24

(2) INFORMATION FOR SEQ ID NO:156:

AAGGATCCGA ACGCTATCTT CCTG

(i) SEQUENCE CHARACTERISTICS:

(A) LENCTH: 56 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:156:

GGGAGCICGI ICAGGGICAG AACCAGAGAG GIACGAGACG GAACAGCAGI GGIIGG

28

(2) INPORMATION FOR SEQ ID NO:157:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 157 base pairs
(B) TYPE: nucleic acid
(C) 'STRANDEDNESS: double
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(*i) SEQUENCE DESCRIPTION: SEQ ID NO:157:

G GAI CCG AAC GCI AIC ITC CTG ICT IIC CAG CAC CTG CTG CGI GGC Asp Pro Asn Ala Ile Phe Leu Ser Phe Gln His Leu Leu Arg Gly 15

Leu G AAA GTT COT ITC CIG AIG CIG GIT GGC GGI ICI ACC CIG IGC GIT CGI Lys Val Arg Phe Leu Het Leu Val Gly Gly Ser Ihr Leu Cys Val Arg 25 CCG CCA ACC ACT GCT GTT CCG TCT CGT ACC TCT CTG GTT Pro Pro The The Ala Val Pro Ser Arg The Ser Leu Val 35 7 8 66 F 66 85

142

157

CTG AAC GAG CTC Leu Asn Glu Leu 50 Acc

(2) INFORMATION FOR SEQ ID NO:158:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 100 base p (B) TYPE: nuclaic acid (C) STRANDEDNESS: sing (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

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CCTCGAGGAA ITCCTGCAGC CCGGGACTAG IATCGGCTAC CCCTACGACG ICCCCGACTA

GCCGGCGTC CATCACCATC ACCATCACTG AGGGGCGGCC

(2) INFORMATION FOR SEQ ID NO:159:

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:158:

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AAITGGGGGC CGCTCAGTGA TGGTGATGGT GATGAGGGCC GGGGTAGTCG GGGACGTCGT

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:159:

(ii) MOLECULE TYPE: Other nucleic acid

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 108 base pairs (B) TYPE nuclaic acid (C) STRANDENESS: single (D) TOPOLOGY: linear

AGGGTAGCC GATACTAGTC COGGCTGCA GGAATTCCTC GAGGGTAC

(2) INFORMATION FOR SEQ ID NO: 160:

(2) INFORMATION FOR SEQ ID NO:163:	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 35 base pairs (B) TYEN inclaid acid (C) STRANDENESS: aingle (D) TOPOLOGY: linear	(ii) MOLECULE TYPE: Other nucleic acid	(x1) SEQUENCE DESCRIPTION: SEQ ID NO:163:	TCAAGCTTAC TAGTCCCTTC CTGAGACAGA TTCTG	(2) INFORMATION FOR SEG ID NO:164:	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 22 base pairs (B) TYPE: nucleic acid (C) STRANDENESS: single (D) TOPOLOGY: linear	(11) MOLECULE TYPE: Other nucleic acid	(x1) SEQUENCE DESCRIPTION: SEQ ID NO:164:	TAATACGACT CACTATAGGG CG	(2) INPORMATION FOR SEQ ID NO:165:	(1) SEQUENCE CHARACTERISTICS: (A) LEWGTH: 22 base pairs (B) TYES nucleic acid (C) STRANDENESS: single (D) TOPOLOGY: linear	(ii) MOLECULE TYPE: Other nucleic acid	(*1) SEQUENCE DESCRIPTION: SEQ ID NO:165:	AATTCCAAGA TCACACTT GC	(2) INPORMATION FOR SEQ ID NO:166:	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 15 base pairs (B) TYBE: nucleic acid (C) STRANDENESS: single (D) TOPOLOCY: linear	(ii) MOLECULE TYPE: Other nucleic acid	(*1) SEQUENCE DESCRIPTION: SEQ ID NO:166:	TCAAGCITAC TAGICCCTIC CIGAGACAGA IICIG	(2) INFORMATION FOR SEQ ID NO:167:	(i) SEQUENCE CHARACTERISTICS: (A) IMPORT: 15 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOCY: linear	(ii) MOLECULE TYPE: Other nucleic acid	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:167:

AATICCAAGA ICTCACACIT GCITITGACA CAACIGIGIT IACITGCAAT CCCCCAAAAC 60

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:160:

(ii) MOLECULE TYPE: Other nucleic acid

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 70 base pairs
(B) TYPE: nucleic acid
(C) STRANDENESS: single
(D) TOPOLOGY: lines

22

35

GGGICTGICI GIIIIGGGGG AITGCAAGIA AACACAGITG IGICAAAAGC AAGIGIGAGA 60

(x1) SEQUENCE DESCRIPTION: SEQ ID NO:161:

(ii) MOLECULE TYPE: Other nucleic acid

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 66 base pairs
(B) TYPE: nuclaic acid
(C) STRANDENESS: single
(D) TOPOLOGY: linear

(2) INFORMATION FOR SEQ ID NO:161:

AGACAGACCC

53

(*i) SEQUENCE DESCRIPTION: SEQ ID NO:162: (ii) MOLECULE TYPE: Other nucleic acid.

GGCCGGGAI GGAGCIGACT GAATTGCTC

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 29 base pairs
(B) TYPE: nucleic acid
(C) STRANDDRESS: single
(D) TOPOLOGY: lines

(2) INFORMATION FOR SEQ ID NO:162:

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(1) SEQUENCE CHANACTERISTICS:
(A) LENGTH: 27 base pairs
(B) TYPE: nucleic acid
(C) STRANDENESS: single
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:168:

TGTAGGCAAA GGGGTAACCT CTGGGCA

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(2) INFORMATION FOR SEQ ID NO:169:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 21 amino acids
(B) TTPE: amino acid
(C) STRANDEDNESS:
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: peptide

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:169:

The Ser Ile Gly Tyr Pro Tyr Asp Val Pro Asp Tyr Ala Gly Val His $\frac{1}{5}$

Xaa Xaa Xaa Xaa Xaa

(2) INFORMATION FOR SEQ ID NO:170:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 3D base pairs
(B) TYPE: nucleic acid
(C) STRANDENESS: single
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:170:

TITGAAITCG GCCAGCCAGA CACCCCGGCC

30

(2) INFORMATION FOR SEQ ID NO:171: (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 41 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(*!) SEQUENCE DESCRIPTION: SEQ ID NO:171:

(2) INFORMATION FOR SEQ ID NO:172:

ITTGCGGCCG CTCAITAITC GIGIAICCIG ITCAGGIAIC C

4

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 24 base pairs
(B) TYPE: nucleic acid

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(C) STRANDEDNESS: single (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(x1) SEQUENCE DESCRIPTION: SEQ ID NO:172:

TGTAGGCAAA GGAACCTCTG GGCA

24

(2) INFORMATION FOR SEQ ID NO:173:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 27 base pairs
(B) TYPE: nucleic acid
(C) SYRANDENESS: aingle
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:173:

TGTAGGCAAA GGAGTGTGAA CCTCTGG

(2) INFORMATION FOR SEQ ID NO:174:

(1) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 27 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(*i) SEQUENCE DESCRIPTION: SEQ ID NO:174:

TGTAGGCAAA GGAGCGTGAA CCTCTGG

27

(2) INFORMATION FOR SEQ ID NO:175:

(A) LENGTH: 27 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (1) SEQUENCE CHARACTERISTICS

(ii) MOLECULE TYPE: Other nucleic acid

(*1) SEQUENCE DESCRIPTION: SEQ ID NO:175:

TCTAGGCAAA GGTCCGTGAA CCTCTGG

27

(2) INFORMATION POR SEQ ID NO:176:

(A) LENGTH: 24 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (1) SEQUENCE CHARACTERISTICS:

(ii) MOLECULE TYPE: Other nucleic acid

(*I) SEQUENCE DESCRIPTION: SEQ ID NO:176:

AGCTGTGGTC CTCTGTGGAG GAAG

(2) INFORMATION FOR SEQ ID NO:177;

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(A) LENGTH: 24 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: single
(D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:177: (ii) MOLECULE TYPE: Other nucleic acid

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(2) INFORMATION FOR SEQ ID NO:178:

24

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 27 base pairs
(B) TYPE: nucleic acid
(C) STRANDENESS: single
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:178:

AGCTGTGGTC CTGTTGCCCT GTGGAGG

(2) INFORMATION FOR SEQ ID NO:179:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 27 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: single

(C) STRANDEDNESS: Bir (D) TOPOLOGY: linear

(11) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:179:

ACCTGTGGTC CTAGGGCCCT GTGGAGG

(2) INFORMATION FOR SEQ ID NO:180:

(1) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 27 base pairs
(B) TYPE: nucleic acid
(C) STRANDENESS: single
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:180:

AGCTGTGGTC CTTCCGCCCT GTGGAGG

27

(2) INFORMATION FOR SEQ ID NO:181:

(1) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 26 base pairs
(B) TYPE: nuclaic acid
(C) STRANDENESS: single
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:181:

CGGCTGCAG GATATCCAAG ATCTCA

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(2) INFORMATION FOR SEQ ID NO:182:

(1) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 22 base pair: (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear

(11) MOLECULE TYPE: Other nucleic acid

(*1) SEQUENCE DESCRIPTION: SEQ ID NO:182:

TAATACGACT CACTATAGGG CG

(2) INFORMATION FOR SEQ ID NO:183:

(1) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 34 base pairs
(B) TYPE: nuclaic acid
(C) STRANDEDNESS: anngle
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: Other nucleic acid

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:183:

(2) INFORMATION FOR SEQ ID NO:184:

GGGGGGCGC TCAGCTGGG ACAGCTGTGG TGG

(A) LENGTH: 7 amino acids
(B) TYPE: amino acid
(C) STRANDEDNESS:
(D) TOPOLOGY: linear (i) SEQUENCE CHARACTERISTICS:

(ii) MOLECULE TYPE: peptide

(ix) FEATURE:

(A) NAME/KEY: misc feature (D) OTHER INFORMATĪON: /note= "The amino acid at position 2 is Ala, Ser, Gly, Het or Gln." .

(A) NAME/KEY: misc feature (D) OTHER INFORMATION: (Lx) FEATURE:

/notes "The amino acid at position 7 is Glu or Lys."

(*1) SEQUENCE DESCRIPTION: SEQ ID NO:184:

Lys Xaa Tyr Tyr Glu Ser Xaa

(2) INFORMATION FOR SEQ ID NO:185:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 6 amino aci
(B) TYPE: amino acid
(C) STRANDENESS:
(D) TOPOLOGY: linear

(11) MOLECULE TYPE: peptide

(ix) FEATURE:
 (A) NAME/KEY: misc_feature
 (D) OTHER INFORMATION:

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/note= "The amino acid at position 6 is Glu or Asp."

(*i) SEQUENCE DESCRIPTION: SEQ ID NO:185:

Lys Glx Arg Ala Ala Xaa

(2) INFORMATION FOR SEQ ID NO:186:

(1) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 7 amino acids
(B) TYPE: amino acids
(C) STRANDENESS:
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: peptide

(*1) SEQUENCE DESCRIPTION: SEQ ID NO:186:

Lys Ala Gly Xaa Cys Ser Gly

(2) INPORMATION FOR SEQ ID NO:187:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 13 amino acids
(B) TYPE: amino acid
(C) STRANDENESS:
(D) TOPOLOCY: linear

(ii) MOLECULE TYPE: peptide

(1x) FEATURE:

(A) NAME/KEY: misc feature (D) OTHER INFORMATION: /note= "The amino acid at position 2 is Ile, Thr or Ser."

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:187:

Lys Xaa Pro Val Pro Pro Ala Cys Asp Pro Arg Leu Leu 1

(2) INFORMATION FOR SEQ ID NO:188:

(1) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 9 amino acids
(B) TYPE: amino acid
(C) STRANDEDNESS:
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: peptide

(*i) SEQUENCE DESCRIPTION: SEQ ID NO:188:

Lys Asp Ser Phe Leu Ala Asp Val Lys

(2) INFORMATION FOR SEQ ID NO:189;

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 9 amino acids
(B) TYPE: amino acid
(C) STRANDEDNESS:

332

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: peptide

(ix) FEATURE:

TAINGE.

(A) NAME/KEY: misc feature

(D) OTHER INFORMATION:

/note= "The amino aid at position 1 is
Lysine or Arginine."

(x1) SEQUENCE DESCRIPTION: SEQ ID NO:189:

Xaa Thr Leu Pro Thr Xaa Ala Val Pro

(2) INFORMATION FOR SEQ ID NO:190:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 15 amino acids
(B) TYPE: amino acid
(C) STRANDEDNESS:
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: peptide

(x1) SEQUENCE DESCRIPTION: SEQ ID NO:190:

Lys Asp Ser Phe Leu Ala Asp Val Lys Gln Tyr Tyr Glu Ser Glu 1 5

(2) INFORMATION FOR SEQ ID NO: 191:

(i) SEQUENCE CHARACTERISTICS:
(A) LEWRIH: 332 amino acids
(B) TOPOLOGY: linear
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: protein

(vi) ORIGINAL SOURCE:
(A) ORCANISM: Homo Sapiens

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:191:

Ser Pro Ala Pro Pro Ala Cys Asp Leu Arg Val Leu Ser Lys Leu Leu 1 5 15

Arg Asp Ser His Val Leu His Ser Arg Leu Ser Gln Cys Pro Glu Val 25

His Pro Leu Pro Thr Pro Val Leu Leu Pro Ala Val Asp Phe Ser Leu 45

Gly Glu Trp Lys Thr Gln Met Glu Glu Thr Lys Ala Gln Asp Ile Leu 50

Gly Ala Val Thr Leu Leu Leu Glu Gly Val Het Ala Ala Arg Gly Gln 65 75 80

Leu Gly Pro Thr Cys Leu Ser Ser Leu Leu Gly Gln Leu ser Gly Gln $90\,$

Val Arg Leu Leu Cly Ala Leu Gln Ser Leu Leu Gly Thr Gln Leu 110

Pro Pro Gin Gly Arg Thr Thr Ala His Lys Asp Pro Asn Ala Ile Phe 125

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CGC Arg 245

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ACC TCT (Thr Ser 1 2 66 147 195 Leu Pro Asn 175 Cys Val Arg Arg Ala Pro Pro Thr Thr Ala 155 Leu Leu Glu Thr Aan Phe Thr Ala Ser Ala Arg Thr 190 Gly Leu Leu Lys Trp Gln Gln Gly Phe Arg Ala Lys Ile 200 Ser Leu Asp Gln Ile Pro Gly 220 2 g g Thr Ser Asp Thr Gly Ser Leu Pro Pro Asn Leu Gln Pro Gly Tyr Ser 265 Pro Thr His Pro Pro Thr Cly Gln Tyr Thr Leu Phe Pro Leu 275 Pro Ser Ale Pro Thr Pro Thr Ser Pro Leu Leu Asn Thr 310 Gln Leu His Pro Leu Leu Pro 300 ម្ពីដ CAT His CCT 176 CCT Val Leu Aen Arg Ile His Glu Leu Leu Aen Gly Thr Arg Gly Leu 235 Gly Ala Pro Asp Ile Ser Ser 250 Ala Ala JCC Ser 553 AGC CCG GAC Asp 1563 Thr Leu Asn Glu 170 **P**FG CTG AGC CAG TGC CCA GAG GTT CAC 13c Ser Ser Tyr Thr His Ser Gin Asn Leu Ser Gin Glu Gly 325 S S re d (*1) SEQUENCE DESCRIPTION: SEQ ID NO:192: ម្ពុន្ធ GAG CTG ACT (Glu Leu Thr (-20 832 (1) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 1086 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: double
(D) TOPOLOGY: linear (vi) ORIGINAL SOURCE: (A) ORGANISM: Homo Sapiens 걸 E S (11) MOLECULE TYPE: CDNA to mRNA Le. Leu Leu Asn Gln Thr Ser Arg 215 Pro Thr Leu Pro Thr Pro Val Val 290 Ęż AGT (2) INFORMATION FOR SEQ ID NO:192: Pro Ser Arg Thr Ser Leu Val Pro Gly Pro Ser Arg Arg Thr Leu 245 Arg -5 53 GOCCAGCCAG ACACCCCGGC CAGA ATG (MEL (A GG CTC Val Gly Gly Ser Thr Leu 뱕 ទូ អ្នក EB 53 CAC AGC AGA 180 ម្ពីជំ GAC Ser Ser 195 537 757 Thr Pro 61y 210 Gly Pro Ser ATG Het Ala Ala E Arg 녎 25° A6p 305 ម្ពិស្តិស Val Cit S T C

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but her cut sid sid case case cut cut sac cut int get bro the Pro Val Val Gin Leu His Pro Leu Leu Pro Asp Pro Ser Ala 300	OCA ACC COC ACC COT ACC ACC COT CIT AAC ACA TOC IAC ACC CAC Pro Thr Pro Thr Pro The Ser Pro Lew Lew Asn The Ser Tyr Thr His 310	TCC CAG AAT CTG TCT CAG GAA GGG TAA Ser Gln Asn Leu Ser Gln Glu Gly 325	(2) INFORMATION FOR SEQ ID NO:193:	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 7 amino acida (B) TYPE: amino acid (C) STRANDENESS: single (D) TOPOLOGY: linear	(ii) MOLECULE TYPE: peptide	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:193:	Lys Gln Tyr Tyr Glu Ser Glu 1	(2) INPORMATION FOR SEQ ID NO:194:	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 1663 base pairs (B) TYPE: anchaic acid (C) STRANDEDNESS: double (D) TOPOLOGY: linear	(ii) HOLECULE TYPE: CDNA to mRNA	(vi) ORIGINAL SOURCE:(A) ORGANISH: Rattus norvegicus(H) CELL LINE: McA-RH8994	(*1) SEQUENCE DESCRIPTION: SEQ ID NO:194:	GOTGIACCIG GGTCCTGAAG CCCTTCTTCA CCTGGAIAGA TICCTTGGCC CACCTGICCC	CACCCCACTC TGTGGAGAGG TACAAAAGGT CAAGGCGGTCT CCATGGCCGC AGGAAAGATT	CAGGGAGAG GCCCCACACA GGGAGCCACT GCAGTCAGAC ACCCTGGGCA GA ATC Met -21	GAG CTG ACT GAI TTG CTC CTG GTG GCC ATA CTI CTC CTC ACC GCA AGA Glu Leu Thr Asp Leu Leu Leu Leu Leu Leu Leu Thr Ala Arg -20	CTA ACT CTG TGC AGC CCA GTT CGT GCC GCC TGT GAC CCC AGA CTC CTA Leu Thr Leu Ser Ser Pro Val Pro Pro Ala Cys Asp Pro Arg Leu Leu 1	AAT AAA CTG CTT GGT GAC TGC TTG CAT CAC GGA CTG AGT CAG Asn Lys Leu Leu Arg Asp Ser Tyr Leu Leu His Ser Arg Leu Ser Gin 15	TGT CCT GAC GTC AAC CCT TTG TCT ATC CCT GTC CTG CCT GCT GTC Cys Pro Asp Val Asn Pro Leu Ser Ile Pro Val Leu Leu Ero Ala Val 35

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415	463	511	559	607	663	723	783	843	903	963	1023	1083	1143	1203	1263	1323	1383	1443	1503	1563	1623	1663
GAC TTT ACC CTG GGA GAA TGG AAA ACC CAG ACG GAA CAG AGG CCA App Phe Ser Leu Gly Glu Trp Lys Thr Gln Thr Glu Gln Ser Lys Ala 45	CAG GAC AIT CTA GGG GCA GTG TCC CIT CTA CTG GAG GGG GTG ATG GCA Gln Asp lle Leu Gly Ala Val Ser Leu Leu Leu Glu Gly Val Het Ala 75	GCA GCA GCA CAG ITG GAA CCC ICC TCC TCC TCC CTC GGA CAG Ala Arg Gly Gln Leu Glu Pro Ser Cys Leu Ser Ser Leu Leu Gly Gln 80 85 90	CTT ICT GCT CAG GTT CGC CTC CTC TTG GGA GCC CTG CAG GGC CTC CTA Leu Ser Gly Gln Val Arg Leu Leu Leu Gly Ala Leu Gln Gly Leu Leu 105	GGA ACC CAG GTA AGT CCC CAG ACC TAI AGA AAC TAC CCT CTT ACT CAG Gly Thr Gin Val Ser Pro Gin Thr Tyr Arg Asn Tyr Pro Leu Thr Gin 110	TIC CIC TAAGGACCIG GGAAAAGACA AGGGATICIA GAITCIAGGI GICTICAGIG Phe Leu 125	INIGANAGCT GETCTATACG GAGTGATGCT TCTCAGCCAC AATACCTGGG TGCTGGCAGT	AAAICITICC ACCITAGIGA GAAGAGGCCI GAIAIGIGGG CCAACICACI GGCCICAGGC	CCAICCICTG CCTICAGCTI CCICCACAGG GCAGGACCAC AGCTCACAAG GACCCCAGIG	CCCICIICII GAGCIIGGAA CAACIGCIIC GGGGAAAGGI GCGCIICCIG CIGCIGGIAG	AAGGICCCG CCTCTGIGIC AGAGGARCC ITCCCACCAC AGCTGICCCA AGCAGAACCI	CTCAACTCCT CACACTAAAC AAGTTCCCAA ACAGACTTCT GGATTGTTGG AGACGAACTT	CAGTGITGIA GCCAGAACTG CTGGCCCTGG ACTTCTGAAC AGGCTTCAAG GAITCAGAGC	CAAGATTAIT CCTCGTCAGC TAAAICAAAC CTCCGGGTCC TTAGACCAAA TCCCTGGATA	CCTGAACGGG ACACACGAAC CTGTGAATGG AACTCATGGG CTCTTTGCTG GGACCTCACT	ACAGACCCTG GAAGCCCCAG ACGTTGTGCC AGGAGCTTTC AACAAAGGCT CCTTGCCACT	CAACCICCAG AGIGGACTIC CICCIAICCC AAGCCIIGCI GCIGAIGGAI ACACACIIII	CCTCCTICA CCTACCTICC CCACCCTGG GICTCCACCC CAGCTCCCCC CCGITICCTG	ACCCTCCAC CACCATACCT AACTCTACCA ACCCTCATCC AGGACTTGGT CTCAGTAAGC	GICCOGIGCA CIGGCAGGA GCGGAICGI CIGCAACAIC ICICAGGGGC AAGCIICCIC	AGGAAGGCTC TGAGGCAGCT CACTAGACAT CCTGCTCTCG CCTAACGGGC CCTGGGAAAG	GGATACACAG GCCAGGACAC TGTACAACCT TAGGAGCGAT ITITITCTIA ACCTAICAAC	aataticatc agaggaaaa aaaaaaaaa aaaaaaaa

(2) INFORMATION FOR SEQ ID NO:195:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 861 base pairs (B) TTPE: nucleic acid (C) STRANDEDNESS: duble (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA to mana

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PCT/JP95/00208 Asn Gin Thr Ser Arg Ser Leu Asp Gin Ile Pro Gly Tyr Leu Asn Arg 215 CAT His Lys A L P S Pro Cr ğ 5538 95.5 9 5 19 CTC GTC Leu Val Ser GAA TGG A Glu Trp I 50 CTC TTARAAAAA AAAAAAAAA AAAAAAAA AAAAAAAAA Leu Ala TCC Ser Tig. GTG Val S S CTC Leu CAG Gln 115 캶 S 5 CAC CCT 1 His Pro 1 AGC CCG Ser Pro CGT GAC GCA Ala 66.3 G Ly Arg F S AGC Ser 130 g g Lea 62. 61.9 65 ខ្លួត re d GTC 5 2 r g CTA Val 11G 7CC Ser GAG CTT ž Ž EB 14 Z 528 S C TTC Phe EB St. 12 re d (xi) SEQUENCE DESCRIPTION: SEQ ID NO:196: CAG GAC ATT Gln Asp Ile AAA CTG Lys Leu 15 23 Ser 617 ATC 61.7 95.2 S 12 ATC GCCCAGCCAG ACACCCCGG CAGA ATG GAG CTG ACT . Het Glu Leu Thr -21 -20 33 арабарара алабарарар аварарара барарара TGC CCA G Cys Pro G 30 (i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 1267 base pairs (B) TYPEs nucleic acid (C) STRANDENESS: double (D) TOPOLOGY: linear 8 5 Ala Ala Fr ad 253 7 th 1ct Ser 당 (A) ORGANISM: Homo Sapiens (F) TISSUE TYPE: liver (ii) MOLECULE TYPE: CDNA to mRNA CCA AGG CTA I AAT Asn 125 GAC ABP 45 GCA Ala AGT 15 TH 663 617 돐 (2) INFORMATION FOR SEQ ID NO:196: AAG GCA C Val A Se CTC TE ភូ ភូ SPS GPs Ęŝ A CG1 ATC Met GTC AGC A Pa 95 S E G GAT Asp GTC Val ORIGINAL SOURCE: GAG ACC Glu Thr GTG Val 뀱 8 5 5 E 3 7 2 5 3 8 AGC Ç Z AAG Lys CCT GCT TGT GAC CTC AGA 25 r ca G C J CAG Gln 105 r g ដូដូ CAC His 9 5 9 5 53 AGC 동결승 GAG G1° Chc Chc TCC Ser 23 8 8 8 Ala 120 CAC GAA C His Glu I 230 CAG ATG G Gln Het G 55 GTC CAC A 200 £39 Ser Ser 23 £ 3 S H (vi) ATG Met Ęŝ ម្ល 2538 re G Pr Th 23 613 ATA Vel The T 700 0,78 85 GTC i k F 3 CAC £3 2 6 2 66 675 723 17.1 147 195 243 435 483 627 291 339 531 579 387 CAT His 5 S Lys Lys 950 9.4 8 2 TCC AGA 935 13 CTG GAC CAA ATC CCC GGA TAC CTG AAC AGG 924 180 180 ដ្ឋដូ 캶 ặ얼 533 GTC 13 gr TTC Phe CTC Ala Ala TCC 176 135 GTG 200 553 666 Ser TCT Sar 195 re d AGC CCG C 89.38 8.38 Ser GAC អូ មួ AL a GGA GLY 6 5 6 5 CCC Pro Act 61y Arg Pro CC 0.5 GAC Bis 66A 659 613 65 GTC CCT Pro 23 GTC Val Arg Arg Pro Cr Arg Gr ខ្លួន GTA Val 145 퉑 GRA TTC Glu Leu : AAT GCC ATC TTC C ASn Ala Ile Phe L 125 GGA CAG G Gly Gln V 95 23 AAC ATT 11e TCC Ser i d **5**58 Ħ Ala 160 Ęŝ Val Val ặ rg 3 (*i) SEQUENCE DESCRIPTION: SEQ ID NO:195: GAC TIT AGC 1 Amp Phe Ser L GPG GLn Pro 175 5 g s Kg Kg Z Z Z GAG GLu ATT 65 g ATC Wet 점심 ខ្លួ Th' 337 95 म है Lya R ម្លីដូខ S G Ser i i i i £ 3 걸 ខ្លួ Ala Ala 3248 GCCCAGCCAG ACACCCCGG CAGA ATG GAG CTG 1 Het Glu Leu 1 -21 -20 (A) ORGANISM: Homo Sapiens (P) TISSUE TYPE: liver 6 5 5 5 5 5 AGA Arg 205 £3 ပို့ မို့ 25 25 g ŧŝ Ph e 2 2 2 3 3 g g Ser AGT GCA AGG CAALA ALG I GCT GTG G ATG GCA G Met Ala A AAG GAT CCC 1 Lys Asp Pro 1 50.00 5 5 E3 Arg 140 P CA P E 200 A 다음 A 4 8 553 818 155 53 GTC Acc AAG Lys 95.5 53 GTC 밝 ORIGINAL SOURCE: E G Val. AGC AAG Lys 245 FP H 95 ដ ម្ពុជ្ញ 238 8 £ 발 Arg CG A Th 61.7 61.7 AGG CAC His Acc Arg 53 ABO 185 95 G ក្តិខ្មុ AGA Arg re. GAG £3. CAC CAC £ 53 reg Ser GAG Glu Ala 120 GAG GLu A GG GTC Val 33 Agp Gac 83 21,00 23 533 Ser Val a H WO 95/21919 Ä GCC 135 135 CAG Glu Lys Lys G CAC 2003 걸 TGC Cys ÉB 43 CTC ATG Met 55 Se ig 539 (vi) g G rer Cd 9 9 9 9 a P 253 5535 Ser 23 23 3 Ala Ala 13 C 535 ATC Het EB F S ĘЗ Pro Pro E G 걸었 Thr 165 E3 CTC Val i kg Ęŝ 5 7 7 8 8 8 8 23 A 7 CH Z 草草

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	AGA Arg	GGA 180	666 61y	CTG	AGG	TCA	ACA Thr 260	Acc Thr	TTC	GCT	CAC His	AGACACTGCC GACATCAGCA	GCCCCCCTG GGAGACAACT GGACAAGATT	TGAAAAGGGA	
	AGC	Ser	177 Ser 195	CTC Leu	AAC	7.00 P.70	GAC	CCA Pro 275	Thr	TCT Ser	Thr	ACAT	SGAC	IGA	
	7. 1.	P P	61y	667 61y 210	<u>F</u> 3	61.4 61.4	Ser	Ser	290 290	E S	Tyr	ខ្ល	ţ		
:	GTC Val	AGG	Thr.	Pro	13C 17t 225	ព្	Thr	F 25	Pro CG	GAC ABP 305	TCC Ser	ğ	Ş	560	
	A 201	ABG	참	II e	GGA Gly	F 845	GGA GLY	Ser	53	7 5	ACA 320	GAC.	200	TACACAGGAC	
	a t	CCA Pro 175	AGA	AAG Lys	Pro CC	CTC	TCA Ser 255	Tat	rs er	ĘĘ	ABn	e P	9	69	
	Thr	553	GCC Ala 190	Ala	ATC Ile	GLY GLY	Ser	213 270	TTC	E 3	£ 3	TAAGGTTCTC	S	GTAAAAGGGA	a
	55	GAG G1u	Ser Ser	AGA Arg 205	5 G	Arg	116	5 5	CTC Leu 285	720	re ra	TAN.			AGAA
	S S	ABn	SCC Ala	TTC Phe	6AC ABP 220	147	GAC	555	Thr.	CAC His 300	P. C.	666 614	TTCCCTGCAG	CAAAGCCCTG	Ĕ
	600 Ala 155	E 3	Thr.	GLy GLy	re C	GGA C1y 235	7 CG	CTC Lea	TAT	Se CT	AGC Ser 315	S S	8	A A GC	ATAAACCTTC
	8 g	246	Phe Phe	g g	Jcc Ser	ABI	666 Ala 250	Agn	6 La 6 La	Glo	Acc	330 330			
	ACG Arg	53	AAC ABn 185	g cac	AGC Arg	Leu	GLy GLy	265 265	6 g .	GTC Val	S S	Ser	TACAGCTCCC	TCCTGAAACC	ACTGTACATT
	GTC Val	GTC	S H	213	JCC Ser	<u> </u>	ÉZ	S i	38 Thr	GTC Val	ACC Thr	re ca	ACAG	Ę,	CTCT
	75. 6.73	Lee Cr	Gre Glu	AAG Lys	ACC Thr 215	GAA GLu	Thr.	25	P 25	295 295	720	ABI			
	120 0	Ser	Leu	5 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3	3 5	CAC His 230	Arg Arg	Ser	ro cr	Thr.	J. Thr	GIn GIn	TTGTCTCATG	ACTT	E
	Thr.	ACC Thr 165	£ 3	re d	Asn	ATA IIe	245 245	91.4 91.4	ELS .	55 6	S C	TCC Ser 325	TTCT	TCCTACTITC	ATCATITIC

⁽²⁾ INFORMATION FOR SEQ ID NO:197:

(ii) MOLECULE TYPE: synthetic DNA

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	84	96	144	192	240	288	336	384	432	480	528	549
	GTT Val	TCC Ser	GCT	aaa Lys	ATG Met 75	66C 61y	CTC	GAT ABP	orc val	GCC Ala 155	CTC	
	Arg 10	5 2	Pro	Thr	Val	£38	Ser	AAG	AAG	Arg Arg	Thr 170	
	CTT T	Arg 25	53	GAG	61.4 61.4	CTC Leu	CAG Gln 105	His	61.7 C1.7	AGG	Leu	
	GAC	Ser	5 3 6	S In	GAN	TCC Ser	Le I	GCT Ala	Arg	GTC	GTC	
	101 Cys	CAC His	GTT Val	ATG Met 55	re c	TCT	Ala Co	THE THE	CTC Leu 135	TGC Cys	CTA Len	
1D NO:197	Ala	CTG Leu	770	5 cl 6	15 J	ren Len	66C 61y	Thr.	re c	CTC Leu 150	Ser	
200	ឡដូ	GTG	Acc	Thr	CTC	TGC Cya 85	Leu Leu	Arg	CAC	Thr	Acc Thr 165	
SEQ 1	P. C. C.	CAC His	8 5	AAA Lys	Th'	Thr	55.35	617 617	S C C	TCC Ser	AGA	٠
	GCT	Ser	CHG 156 35	17 J	GTA Val	5 5	CTC	CAG Gla 115	TTC Phe	61y	AGC	
SEQUENCE DESCRIPTION:	8 5°	GAC	8 5	55 G. S.	Ala .	61.y	Arg	Pro Pro	AGC Ser 130	61.7	970 Pro	13
SCR	Ser	95 1	CAC His	GLY	65 65 65	£3	GTT Val	Pro Pro	E G	GTA Val 145	GTC Val	
20 21	61.y	E 3	GTT Val	re C	2 3	55.08 8 0.08	5 to 15	CTC	TTC Phe	£3	SCT Ala 160	TAATGAGAAT
UENC	Arg	5 3 2	es a	Ser	ATC 11e	61. 61.4	617 95	S La	AIC Ile	ATC Met	a i	
	8 2	ĘŻ	8558	TT ag	GAC	A. 25.	Ser	Thr	SCC Ala	Cic	Thr	CTC
(xt)	GTT Val	Ser 1	Cy B	GAC ABP 45	G Ln	A la	ie i	61.y	AAT ABD 125	Phe Phe	ទី ទី	GAG Glu
	15 Lat.	150	cac 61n	GTC	Ala 60	N. P. La	CAC Gla	Cer Cer	Pro Pro	Arg 140	Pro Pro	AAC

⁽¹⁾ SEQUENCE CHARACTERISTICS:
(A) LENGTH: 549 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: double
(D) TOPOLOCY: linear

*

What is claimed is:

- 1. A thrombopoletin (TPO) polypeptide having the biological activity of specifically stimulating or increasing platelet production and comprising the amino acid sequence 1 to 332 of SEQ ID NO:6 or derivative thereof.
- The TPO polypeptide derivative according to claim 1 consisting of the amino acid sequence 1 to 163 of SEQ ID NO:6.
- The TPO polypeptide derivative according to claim 1 consisting of the amino acid sequence 1 to 232 of SEQ ID NO:6.
- 4. The TPO polypeptide derivative according to claim 1 consisting of the amino acid sequence 1 to 151 of SEQ ID NO.8
- 5. The TPO polypeptide derivative of claim 1, 2 or 3 having from 1 to 6 amino terminal amino acids deleted.
- 6. The TPO polypeptide derivative according to claim 2 selected from the group consisting of [Δ His³³]TPO (1-163), [Δ Arg¹¹⁷]TPO (1-163) and [Δ Giy¹¹⁶]TPO (1-163).
- 7. The TPO polypeptide derivative according to claim 2 selected from the group consisting of [His³3, Thr³3', Pro³4|TPO (1-163), [His³3, Ala³3', Pro³4|TPO (1-163), [His³3, Gly³3', Pro³4, Ser³8]TPO (1-163), [Gly¹16', Asn¹16', Asn¹17]TPO (1-163), [Gly¹16', Arg¹¹7]TPO (1-163), [Gly¹16', Arg¹¹7]TPO (1-163) and [Ala¹, Val³]TPO (1-163).

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- 8. The TPO polypeptide derivative according to claim 1 selected from the group consisting of [Thr³³, Thr³³³, Ser³³⁴, Ile³³⁵, Gly³³⁶, Tyr³³⁷, Pro³³⁸, Tyr³³⁹, Asp³⁴⁰, Val³⁴¹, Pro³⁴², Asp³⁴³, Tyr³⁴⁴, Ala³⁴⁵, Gly³⁴⁶, Val³⁴⁷, His³⁴⁸, His³⁵⁰, His³⁵¹, His³⁵², His³⁵³]TPO and [Asn²⁵, Lys²³¹, Thr³³³, Ser³³⁴, Ile³³⁵, Gly³³⁶, Tyr³³⁷, Pro³³⁸, Tyr³³⁹, Asp³⁴⁰, Val³⁴¹, Pro³⁴², Asp³⁴³, Tyr³⁴⁴, Ala³⁴⁵, Gly³⁴⁶, Val³⁴⁷, His³⁴⁸, His³⁵⁰, His³⁵⁰, His³⁵¹, His³⁵², His³⁵², His³⁵¹, His³⁵², His³⁵⁰, His³⁵¹,
- 9. The TPO polypeptide derivative according to claim 1 selected from the group consisting of [Asn 25]TPO and [Thr 33]TPO.
- 10. The TPO polypeptide derivative according to claim 2 selected from the group consisting of [Ala¹, Val³, Arg¹²3]TPO (1-163), [Ala¹, Val³, Arg¹³3]TPO (1-163), [Ala¹, Val³, Leu³2]TPO (1-163), [Ala¹, Val³, Leu³2]TPO (1-163), [Ala¹, Val³, Pro¹⁴8]TPO (1-163), [Ala¹, Val³, Pro¹⁴8]TPO (1-163), Arg⁵³]TPO (1-163) and [Ala¹, Val³, Arg³¹5]TPO (1-163).
- The TPO polypeptide derivative according to claim 2 selected from the group consisting of [Arg¹²⁹]TPO (1-

- 12. The TPO polypeptide according to claim 1 covalently bonded to a polymer.
- The TPO polypeptide according to claim 12 wherein said polymer is polyethylene glycol.
- 14. The TPO polypeptide according to any of claims 1 through 13 which further comprises the amino acid $\rm Met^{-2}\text{-}Lys^{-1}.$
- The TPO polypeptide according to any of claims 1 through 13 which further comprises Met⁻¹.
- 16. The TPO polypeptide according to any claims 1 through 13 which further comprises $\mbox{\rm Gly}\mbox{-}1.$
- 17. A polypeptide consisting of the mature sequence of amino acids of SEQ ID NOs:2, 4 or 6.
- 18. A DNA encoding a TPO polypeptide according to any of claims 1 through 11, 14, 15 and 17.

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- 19. A DNA sequence for use in securing expression of a TPO polypeptide having the biological activity of specifically stimulating or increasing platelet production, which is selected from the group consisting of:
- (a) the DNA sequences shown in SEQ ID NOS:194, 195 or 196 or complementary strands thereof; and
- (b) DNA sequences which hybridize under stringent conditions to the DNA sequences as defined in (a) or fragments thereof; and
- (c) DNA sequences which would hybridize to the DNA sequences as defined in (a) and (b), but for the degeneracy of the genetic code.
- 20. A cDNA sequence according to claim 19.
- 21. A genomic DNA sequence according to claim
- . 6
- A manufactured DNA sequence according to claim 19.
- 23. A process for producing a TPO polypeptide comprising the steps of:

- a) expressing a polypeptide encoded by a DNA according to any of claims 18, 19, 20, 21 or 22 in a suitable host, and
- b) isolating said TPO polypeptide.
- 24. The process of claim 23 wherein the TPO polypeptide expressed is a Met⁻²-Lys⁻¹ polypeptide and further including the step of cleaving Met⁻²-Lys⁻¹ from said isolated TPO polypeptide.
- expression of a TPO polypeptide having the biological activity of specifically stimulating or increasing platelet production, said DNA further encoding a thrombin recognition peptide 5' to TPO polypeptide encoding sequences and encoding glutathione-S-transferase (GST) 5' to the thrombin recognition peptide.
- 26. The DNA sequence of claim 25 for use in securing expression of a TPO polypeptide consisting of the amino acid sequence 1 to 174 of SEQ ID NO:6.
- 27. A process for producing a TPO polypeptide having the biological activity of specifically stimulating or increasing platelet production comprising the steps of:

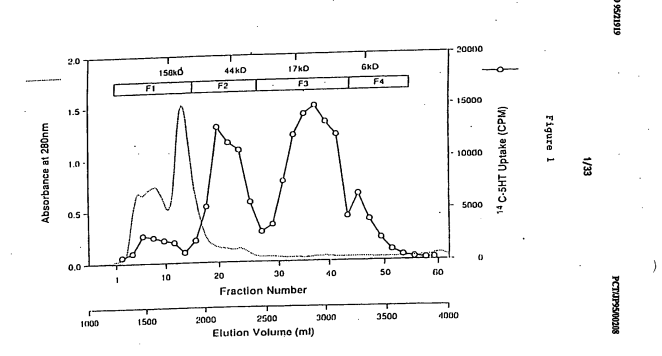
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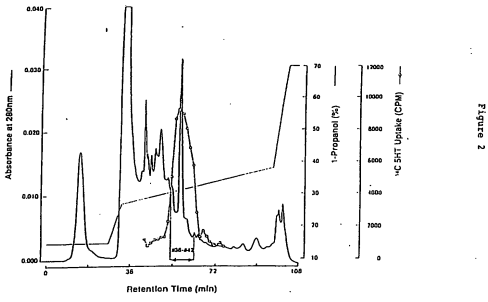
- a) expressing a polypeptide encoded by a DNA according to claim 25 or claim 26 in a suitable host, and
- b) isolating said GST-(thrombin recognition peptide)-TPO polypeptide.
- c) treating said isolated GST-(thrombin recognition peptide)-TPO polypeptide with thrombin, and
- d) isolating a Gly-1-TPO polypeptide.
- 28. The process of claim 27 wherein the Gly-1-TPO polypeptide is [Gly-1]TPO (1-174).
- 29. The process of claim 27 or claim 28 wherein the isolated Gly-1-TPO polypeptide is a TPO derivative consisting of the amino acid sequence 1 to 174 of SEQ ID NO:6.
- 30. A protein product obtained by the process of claims 23, 24, 27, 28 or 29.
- 31. A procaryotic or eucaryotic host cell transformed or transfected with the DNA sequence according to any of claims 18, 19, 20, 21, 22, 25 or 26 in a manner enabling said host cell to express a polypeptide having the biological activity of specifically stimulating or increasing platelet production.

- 32. A pharmaceutical composition comprlsing an effective amount of the polypeptide according to any of claims 1 through 16 and 30 in combination with a pharmaceutically acceptable carrier.
- 33. A pharmaceutical composition according to claim 32 for use in the treatment of platelet disorders.
- 34. A pharmaceutical composition according to claim 32 for use in the treatment of thrombocytopenia.
- 35. A pharmaceutical composition according to claim 34 for use in the treatment of thrombocytopenia induced by chemotherapy, radiotherapy or bone marrow transplantation.
- 36. A method for treating platelet disorders, which comprises administering an effective amount of the polypeptide according to any of claims 1 through 16 and 30 to patients having the disorders.
- 37. A method for treating thrombocytopenia, which comprises administering an effective amount of the polypeptide according to any of claims 1 through 16 and 30 to patients having thrombocytopenia.

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- 38. A method according to claim 37 for treating thrombocytopenia induced by chemotherapy, radiotherapy or bone marrow transplantation.
- 39. An antibody specifically immunoreactive with a polypeptide of any claims 1 through 16 and 30.
- 40. Use of the antibody of claim 39 for the isolation of a polypeptide according to any of claims 1 through 16 and 30.
- 41. Use of the antibody of claim 39 for the quantification of a polypeptide according to any of claims 1 through 16 and 30.

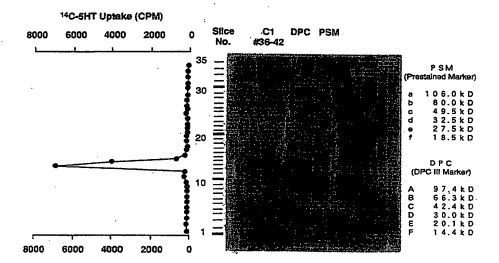


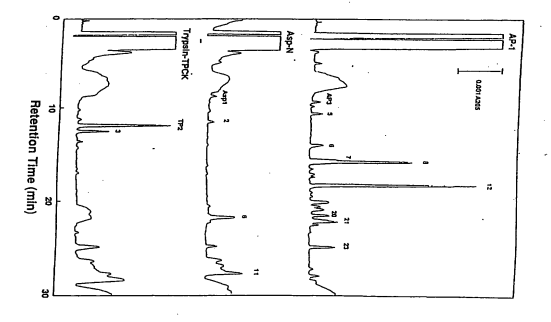


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Figure 3

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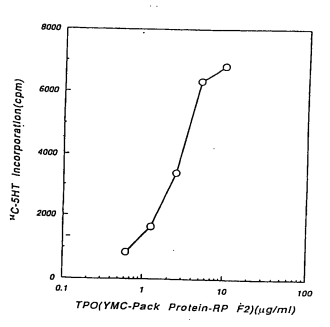
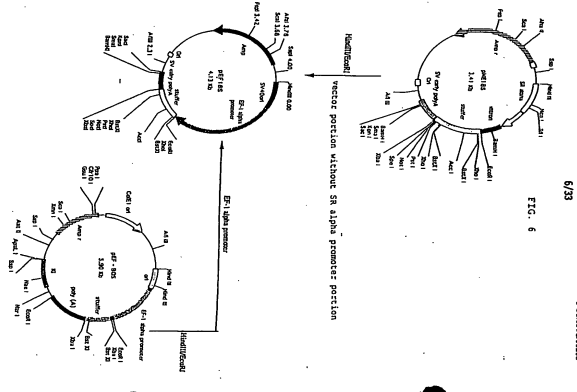
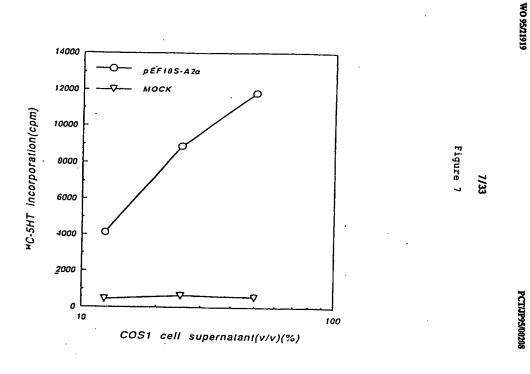


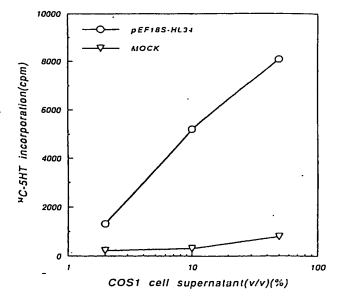
Figure 5

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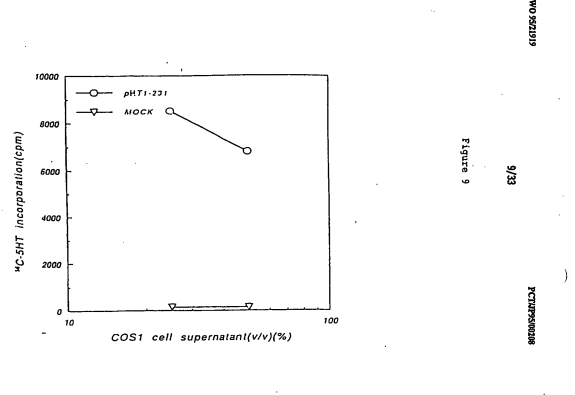


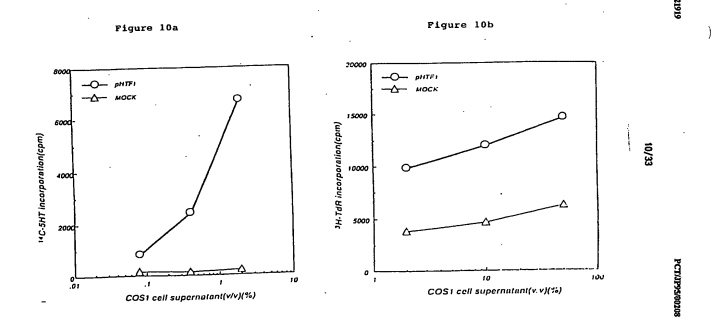


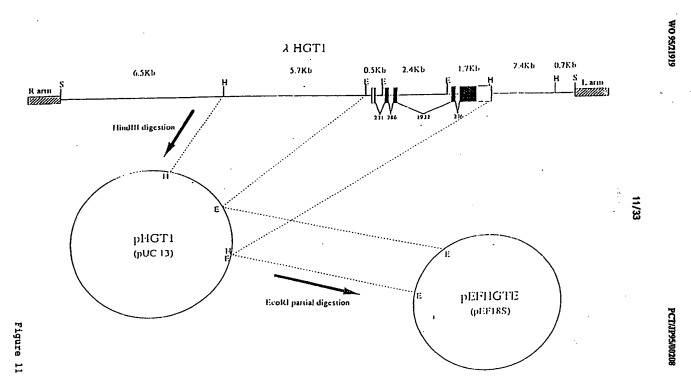
8/33 Figure 8

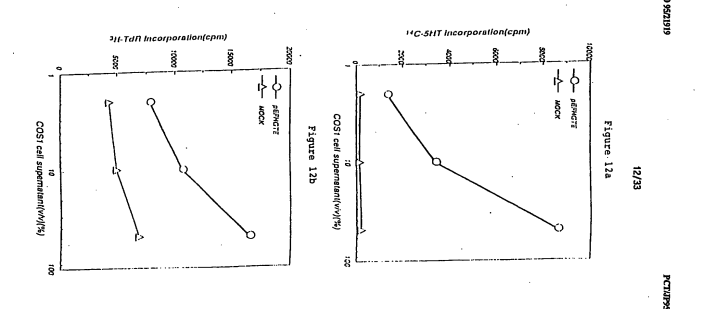
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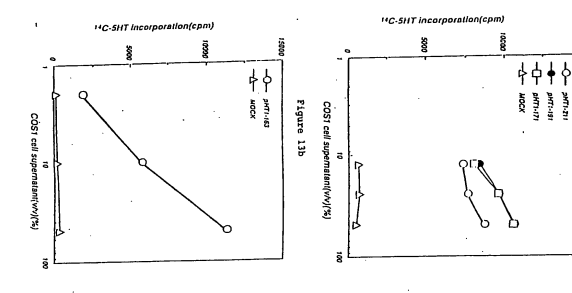


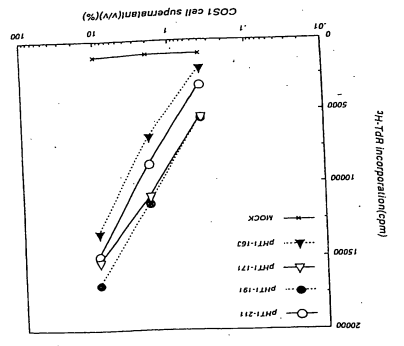


Figure 13a

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Figure 14

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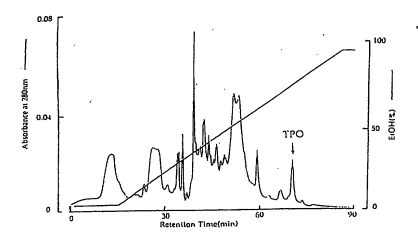
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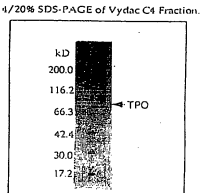


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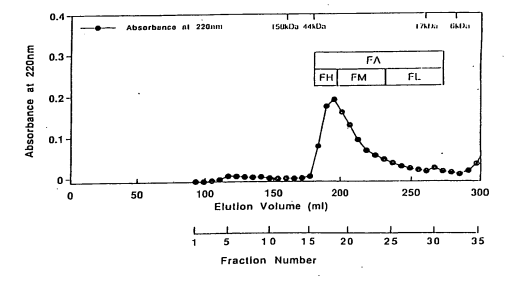


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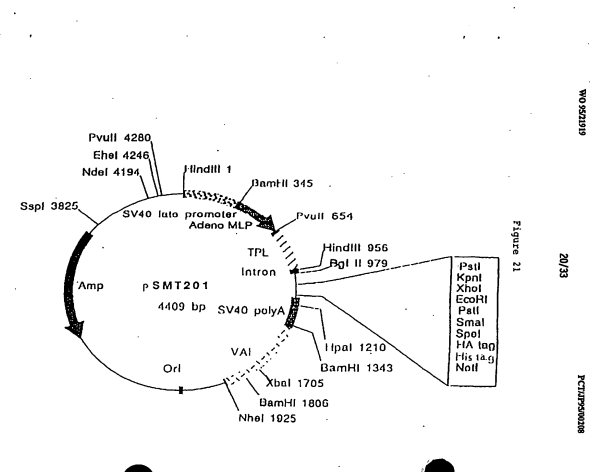


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18/33 Figure 19

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16/25% SDS-PAGE of Putitied hTPO163 (100ng each) < Superdex 75 pg Fractions of Source 15RPC >

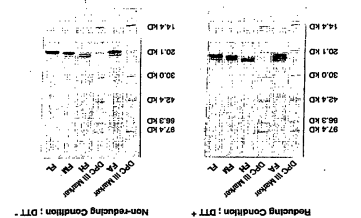
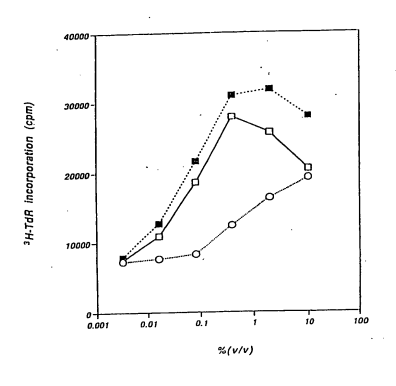
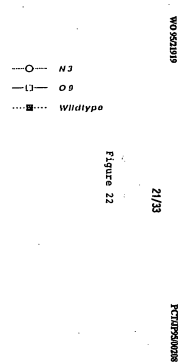
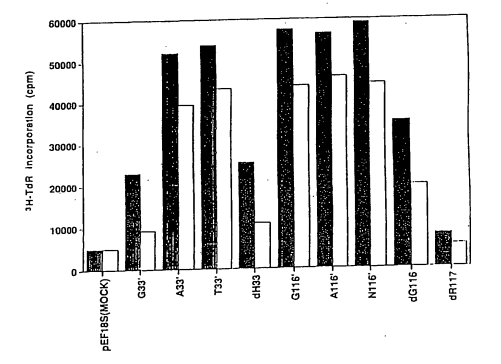


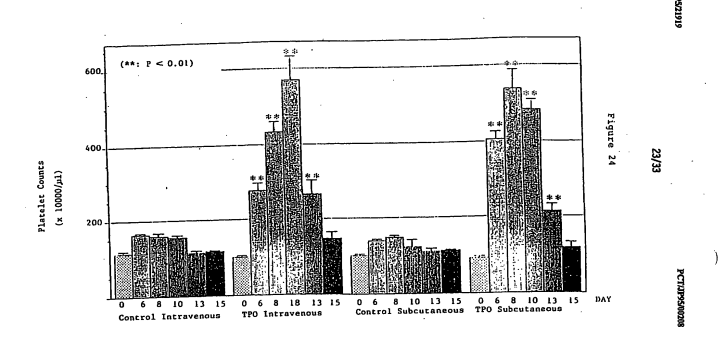
Figure 20







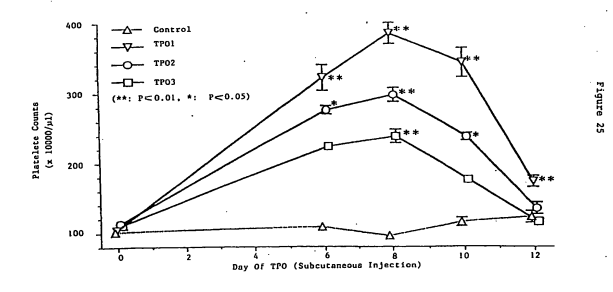


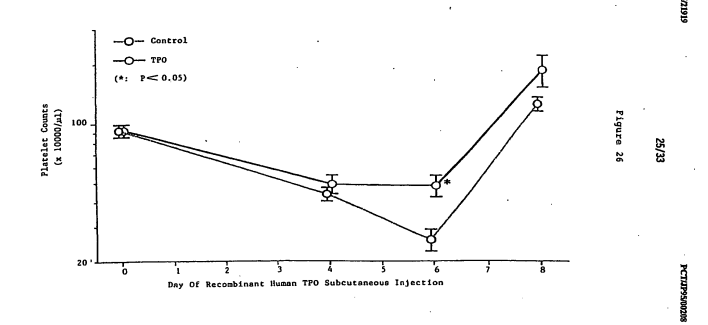


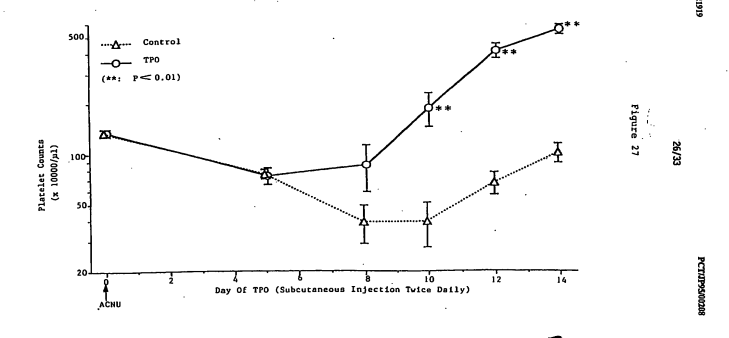
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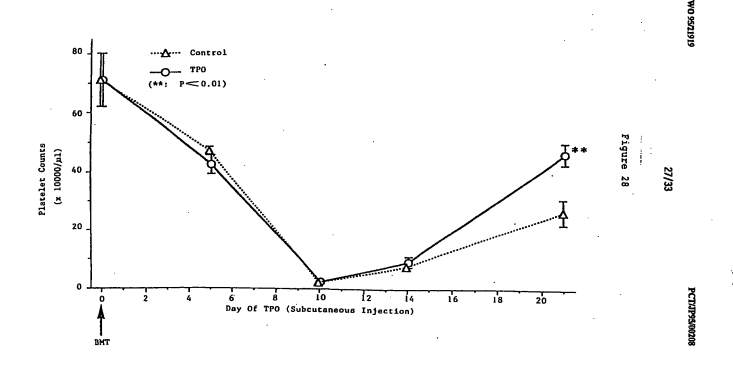
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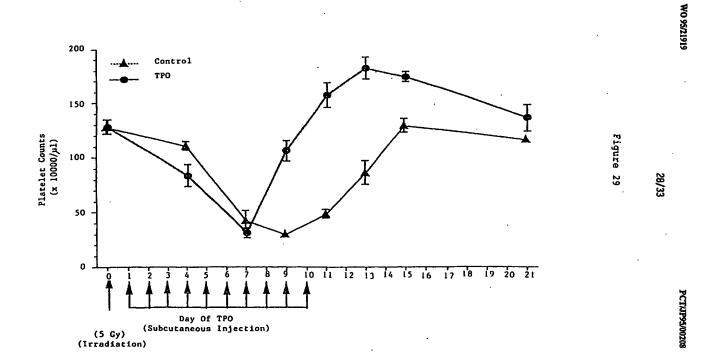
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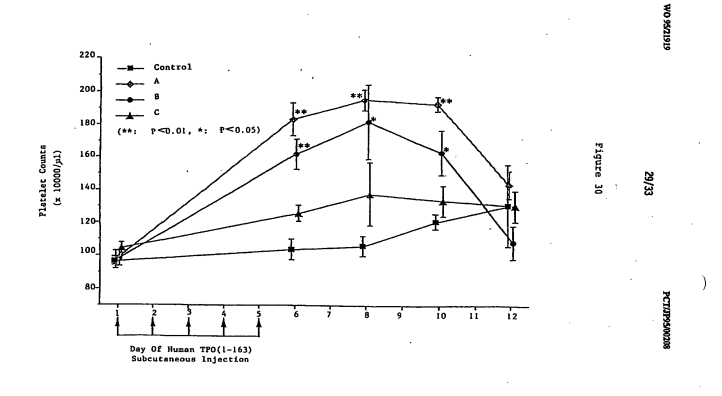












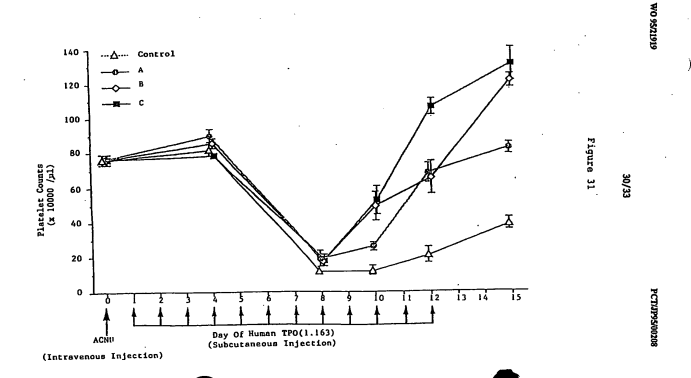
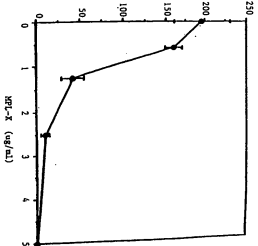


Figure 32



Megakaryocytes per well

NPL-X completely blocks the ability of APK9 to induce megakaryocyte development

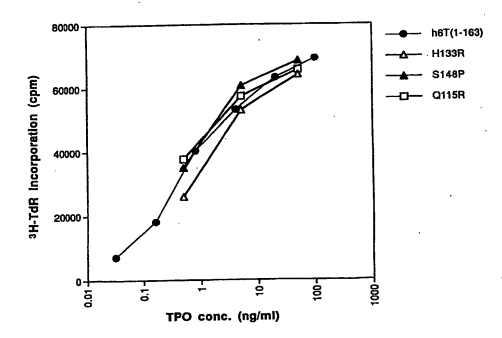
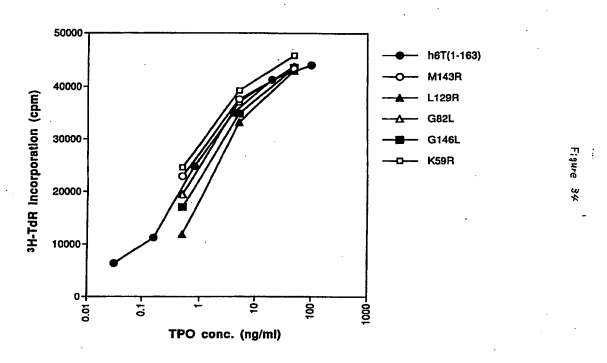


Figure 33

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